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THE Volatility Process : A study of stock market dynamics via parametric stochastic volatility models and a comparison to the information embedded in the options price

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Abstract

This thesis is centered upon developing methodologies for estimation of stochastic volatility parameters from time-series under a classical framework.

The first chapter of the thesis presents a general introduction and a concise survey of various nondeterministic volatility models. Some popular models such as Heston or VGSA are discussed in more detail.

The second and principal chapter lays out our time-series inference methodologies, which are mostly based on the maximization of likelihood. Employing particle-filters, we implement a systematic and flexible algorithm usable for any parametric stochastic volatility model.

Moreover, we study the reliability of these procedures by running them against simulated data. We will examine the choice of the filter, the time-step and the data length. We also will test the fitting performance of various models upon real market-data. Again, we shall focus on a few examples such as Heston, Bates and VGSA.

The third chapter contains a few examples of application of the previously constructed inference tools. The main example involves the comparison of the skewness of the distributions implied from the stock and options markets. We shall observe an excess negative-skewness in the latter, which may imply either a model misspecification or a trade opportunity. A skewness transaction based on this idea will then be described and tested.
Résumé

I

Le sujet de cette thèse est la Volatilité Stochastique. Le fait que la volatilité des actions soit non-déterministe est accepté depuis les années soixante où Samuelson et Mandelbrot ont décrit le caractère aléatoire de la variance de la distribution du marché des actions.

Cet aspect aléatoire de la volatilité a des conséquences immédiatement visibles sur la distribution des rendements. En effet, celle-ci n'est Normale que de manière conditionnelle. La distribution marginale du rendement sera asymétrique et leptokurtique.

Les conséquences existent également sur l'évaluation des prix d'options et autres produits dérivés. La formule bien connue de Black-Scholes-Merton ne sera plus valide une fois que la volatilité devient non-déterministe.

Il est donc essentiel de noter que ce caractère aléatoire est doublement observable. D'une part, la série temporelle du marché des actions nous fournit des rendements qui ne sont pas Normaux ; et d'autre part les volatilités implicites aperçues dans le marché des options, pour une maturité donnée et pour des prix d'exercice différents, sont non-constantes.

Ce dernier phénomène est généralement dénommé le smile ou le skew de la volatilité. Plusieurs théories existent pour expliquer ce smile. La volatilité stochastique est l'une des théories principales. Une autre est l'existence de sauts dans le prix du sous-jacent, comme Robert Merton l’a mentionné dès les années soixante-dix.

Une manière différente d'aborder le problème serait d'utiliser les prix d'option et en déduire la distribution implicite du rendement du prix de l’action. Cette approche nous donne la possibilité de déterminer ce qui est habituellement appelé la volatilité locale. Cette notion a été introduite par Bruno Dupire au début des années quatre-vingt-dix. On peut calculer cette volatilité locale soit de manière analytique, soit par l’intermédiaire d’un arbre implicite comme le proposent d’une part Rubinstein et Jackwerth, et d’autre part Derman et Kani. Ces arbres nous offrent les probabilités risque-neutres et implicites dans les prix d’options. Ayant le choix entre les options Put et Call, on choisit souvent celles qui sont hors-monnaie pour des raisons de liquidité.

Certains, comme Cox et Ross (CEV) ou encore Bensoussan, Crouhy et Galai (BCG) introduisent le concept d’une volatilité qui dépend du niveau du prix de l’action. Les modèles CEV représentent une classe particulière de la volatilité stochastique où le coefficient de la corrélation entre le rendement du sous-jacent et la volatilité instantanée, est égal à 1.0

Le modèle BCG possède l’avantage d’avoir une interprétation financière : En effet, dans ce modèle, similairement à un modèle proposé par Merton, la valeur de l’action est elle-même une option sur la valeur des actifs de l’entreprise. Dans ce cas là, ces actifs, sur lesquels il n’y a pas de marché, ont leur propre volatilité. Or il est possible d’éliminer
cette volatilité des équations et obtenir une relation directe entre l’action et sa volatilité. Cette dernière est, par conséquent, une fonction implicite de l’action.
Quant aux modèles de type CEV, ses auteurs spécifient la relation entre la volatilité et l’action explicitement. Cette dépendance est en fait artificielle, mais elle simplifie les calculs considérablement.

La méthode d’économétrie pour introduire la notion de la volatilité stochastique serait l’utilisation de modèles de type \textit{GARCH}, où le bruit du rendement du prix constitue une série temporelle qui elle-même est composée de variables aléatoires. Ce qui est plus, les termes de cette série sont auto-corrélés.
Cela dit, il est primordial de noter que sous ce modèle, la variance est parfaitement \textit{connue} à une date donnée, comme fonction des rendements historiques de même que des variances précédentes. Ceci est différent de la théorie générale de la volatilité stochastique où l’on a un modèle à deux facteurs. En effet, pour un tel modèle la variance n’est pas parfaitement connue à une date donnée et représente une variable aléatoire. L’idée fondamentale derrière le modèle GARCH, est basée sur le fait qu’en pratique, on observe qu’une volatilité élevée est suivie par une période de volatilités élevées et vis versa. Ce phénomène dénommé \textit{cluster} pourrait en effet être modélisé par un GARCH.

On a dit que pour évaluer les prix d’option, on ne peut utiliser les formules classiques supposant une volatilité constante. Les méthodes disponibles pour le calcul de prix d’options sous volatilité stochastique comprennent plusieurs catégories :

La première catégorie est celle des méthodes de Différences Finies (FD) appliquées aux Equations aux Dérivées Partielles (PDE) Cette méthode nous donnera une grille à trois dimensions : le temps, le prix de l’action et le niveau de la volatilité instantanée. Un des avantages de cette méthode est qu’elle peut évaluer les options américaines et prendre en compte la possibilité de l’exercice prématuré.

La deuxième catégorie est celle de méthodes d’inversion de transformées de Fourier. En effet, pour la plupart des modèles de volatilité stochastique existant dans la littérature, la transformée de Fourier est disponible de manière analytique. Le modèle de Heston est l’exemple le plus connu concernant ce fait. L’inversion de cette transformée pourrait être effectuée numériquement comme l’ont suggéré par exemple Carr et Madan.

La troisième catégorie est celle de modèles à simulation Monte-Carlo. Cela pourrait se faire soit directement en prenant les équations à deux facteurs, soit en utilisant des techniques de \textit{Mixage} de Hull et White ou encore Romano et Touzi. Dans ces méthodes, on simule la variable aléatoire représentant la volatilité instantanée, puis on applique la formule de Black-Scholes à un prix d’action modifié et une volatilité composite correspondant au chemin simulé.

Finalement il existe des méthodes de développement en séries. Par exemple Alan Lewis suggère un développement sur la volatilité de la volatilité. Cette série devient analytiquement concise pour certains modèles populaires tels que celui de Heston. Autres séries utilisent le développement asymptotique pour maturités longues.
Il est intéressant de noter que contrairement au cas de la volatilité constante ou déterministe, sous une volatilité stochastique, les marchés ne sont pas complets. Cela veut dire que l’on ne peut pas dupliquer la valeur d’une option en utilisant le sous-jacent et des obligations. En effet, en plus du risque associé au sous-jacent, il existe maintenant une deuxième source d’incertitude indépendante de la première. Cette deuxième source est le prix de marché du risque associé à la volatilité, qui est encore une fois totalement distincte du prix de marché du risque associé au sous-jacent même. Certains considèrent malgré cela que les marchés sont complets. En effet on peut toujours générer la valeur des produits dérivés si l’on se sert d’une option en plus du sous-jacent et l’obligation.

Une autre catégorie populaire de modèles de volatilité stochastique rejette l’idée de la diffusion complètement et utilise des sauts. Ceci est différent des modèles de Merton ou Bates où l’on superpose des sauts de Poisson à la distribution Gaussienne. Ici on détient un modèle purement basé sur des sauts. Autrement dit, les modèles de Merton et Bates expliquent les mouvements infinitésimaux du sous-jacent à travers une distribution Normale, et les mouvements plus grands et plus brusques par des sauts. En revanche les modèles de sauts purs, expliquent même les mouvements infinitésimaux avec des sauts. L’exemple le plus populaire parmi ces modèles est celui de Variance Gamma (VG) de Madan et Carr, de même que ses modèles dérivés comme VGSA ou VGG. Comme le nom l’indique, la distribution de base utilisée ici est une distribution Gamma appliquée au temps.

Ainsi ces modèles introduisent la notion de la volatilité stochastique grâce à une randomisation du temps. Autrement dit, les mouvements du sous-jacent s’effectuent non pas de manière chronologiquement uniforme, mais de manière aléatoire. Cela crée les propriétés désirées d’une distribution leptokurtique et asymétrique.

Le modèle VGSA ajoute à ceci une randomisation du temps d’arrivée dans la distribution Gamma. Cela permet d’obtenir les fameux clusters de volatilité, observés en pratique dans le marché. L’évaluation des prix d’options sous ces modèles peut encore une fois se faire soit par l’intermédiaire de l’inversion de transformées de Fourier, soit en utilisant des simulations Monte-Carlo.
Nous avons donc commencé par décrire les différents modèles de volatilité stochastique qui existent dans la littérature. Nous abordons par la suite, le problème fondamental du *calibrage* des paramètres pour ces modèles. L’estimation des paramètres peut se faire soit en utilisant des séries temporelles de prix d’actions, soit en utilisant des prix d’options à une date donnée. Naturellement on pourrait également utiliser des séries temporelles de prix d’options.

Le calibrage via les options, est celui qui est habituellement utilisé dans le marché. Il nous permet, en minimisant par exemple la somme des carrés des différences entre les prix de marché et les prix théoriques des options, d’obtenir les paramètres optimaux. Notez que ces paramètres sont obtenus dans l’univers risque-neutre, où les options sont couvertes dynamiquement, et par conséquent le drift du sous-jacent est remplacé par le taux d’intérêt. Comme on l’a déjà signalé, les prix théoriques devront être calculés avec des modèles par exemple à base de simulations Monte-Carlo ou à base d’inversion de transformée de Fourier.

Plusieurs questions se posent à cette occasion. Quelles options doit-on choisir ? Souvent on répond à cette question par des arguments de liquidité. Typiquement les options proches et hors la monnaie sont celles qui sont plus utilisées.

Combien de prix d’exercice doit-on inclure dans le processus de l’optimisation ? Là, il existe un compromis entre la fiabilité des prix (donc encore une fois la liquidité des options) et l’exhaustivité du processus.

La même question se pose pour les maturités. Idéalement, on devrait obtenir les mêmes paramètres pour deux maturités différentes. Autrement dit, ces paramètres devraient être temporellement homogènes. Mais en pratique ceci peut ne pas être le cas, ce qui pourrait indiquer une mauvaise spécification du modèle.

De plus, certains pensent qu’il faut associer des poids aux options utilisées dans l’optimisation. En effet les options plus proche de la monnaie ont plus d’importance que les autres. On pourrait par exemple se servir des taux de changement de ces options relativement à la volatilité, communément dénommés *vega*.

Quant à la méthode d’optimisation, elle peut être choisie de manière relativement libre, comme la fonction sous l’optimisation est typiquement une somme de carrés de différences de prix, et par conséquent continue et différentiable.

Le calibrage via les séries temporelles est un peu plus subtil. En effet la difficulté que l’on rencontre immédiatement est que la volatilité instantanée n’est pas observable dans les marchés.

Pour obtenir les paramètres, on peut par exemple utiliser une méthode de Maximisation de la Vraisemblance (MLE)

Mais précisément parce que l’on a cet état non-observable (latent), on ne peut calculer la vraisemblance directement. D’où la nécessité d’un *Filtre* pour intégrer l’état inobservable.
Il existe deux étapes dans ce procédé: La Prédiction et le Filtrage. Au cours de la Prédiction, on met à jour la distribution a priori de l’état latent au temps $t$ en utilisant toute l’information que l’on a avant cette date. Cela équivaut à résoudre une équation de Fokker-Planck. Au cours du Filtrage, on obtient la distribution a posteriori de l’état en utilisant d’une part la distribution a priori et d’autre part la vraisemblance conditionnelle. On fait ceci en se servant du théorème de Bayes : A un temps $t$ donné, la distribution a posteriori de l’état, est égale à sa distribution a priori, multipliée par la vraisemblance conditionnelle, et divisée par la vraisemblance inconditionnelle. Or pour déterminer cette vraisemblance inconditionnelle (totale) à un temps donné, on a besoin de calculer l’intégrale associée, ce qui n’est pas toujours réalisable.

Un cas particulier fondamental correspond à celui du filtre de Kalman où les équations sont linéaires et les bruits sont Gaussiens. En effet dans ce cas, les distributions à priori et à postérieure sont Normales et par conséquent, peuvent être caractérisées par leurs deux premiers moments. Cela va sans le dire, cette propriété simplifie les calculs considérablement et tout peut se faire de manière analytique.

Or, si l’on suppose que l’équation différentielle stochastique (SDE) du rendement du prix de l’action observable, de même que celle de la volatilité instantanée, sont quasi-linéaires et ne contiennent que des variables aléatoires Normales, alors on pourra utiliser un Filtre de Kalman Étendu (EKF) Dans l’EKF, on effectue un développement de Taylor de premier ordre sur les fonctions non-linéaires de transition et d’observation. Le reste peut se faire identiquement à un filtre de Kalman traditionnel.

Si l’on juge que les non-linéarités sont importantes mais que les distributions restent conditionnellement Normales, alors un Filtre de Kalman Unscented (UKF) ou un Filtre Non-Linéaire de Kushner (NLF) devra être utilisé. Là, on ne fait plus d’approximation linéaire, on utilise les véritables fonctions non-linéaires mais on suppose que les distributions restent plus ou moins Gaussiennes. Cela nous permet de travailler encore une fois avec les deux premiers moments uniquement.

UKF utilise des points spéciaux appelés les points sigma pour calculer ces moments. NLF utilise les racines de polynômes de Gauss-Hermite à cette fin. De plus NLF contient une étape itérative d’alignement de la distribution de la vraisemblance conditionnelle, et de la distribution a priori de l’état. En effet, si ces les deux densités sont concentrées sur des régions différentes, la convergence de l’algorithme peut en être très affectée.

Si maintenant on ne suppose plus que les variables aléatoires sont Normales, même conditionnellement, et que par exemple elles contiennent des sauts de Poisson ou plus généralement correspondent à des distributions de type Lévy quelconque, alors on devra utiliser des Filtres à Particules (PF) où l’on simule des points par leur SDE discrétisée, avec la méthode Monte-Carlo et l’on associe à chacune de ces particules un poids correspondant à la vraisemblance de l’observation conditionnée par l’état simulé.
Ceci est équivalent à calculer l’intégrale de la vraisemblance via une méthode d’échantillonnage d’importance ou encore un changement de numéraire.

Au lieu de simuler directement par la SDE de l’état, on peut également simuler par une distribution plus simple, et ensuite modifier les poids pour compenser cela. Cette étape permet de tenir compte de la vraisemblance dans l’étape simulation. Des exemples de filtres de ce type sont Filtres à Particules Étendus (EPF) ou Filtres à Particules Unscented (UPF)

Une raison importante pour faire cela, est liée à la remarque que l’on a faite à propos de NLF. En effet, lorsque la vraisemblance conditionnelle et la distribution a priori ne sont pas alignées, la convergence de l’algorithme peut être nettement détériorée. C’est pour cette raison que l’on devrait simuler d’une distribution candidate qui contient déjà une certaine information sur l’observation, et par conséquent s’aligne mieux avec la vraisemblance. La correction des poids pourrait par la suite se faire via la même méthode d’échantillonnage d’importance.

Notons que tout ceci était connu depuis longtemps. Cependant ces Filtres à Particules divergeaient souvent, parce que la variance des poids que l’on vient de mentionner, croît à chaque étape. La raison pour laquelle on peut utiliser ces algorithmes, c’est que récemment des techniques de réduction de variance appelées ré-échantillonnage ont été introduites par des chercheurs tels que Neil Gordon, Adrian Smith et Arnaud Doucet.

Ces techniques permettent d’éliminer les particules simulées avec des poids trop petits et reproduire celles avec des poids énormes. Après cela, on remet tous les poids uniformément égaux à l’inverse du nombre initial des particules. Différentes techniques de ré-échantillonnage existent, cependant elles sont toutes basées sur cette même idée de la réduction de la variance des poids.

L’avantage majeur des Filtres à Particules est leur flexibilité. En effet, du moment que l’on sait simuler par la SDE de la volatilité, et que l’on connaît la densité conditionnelle du prix de l’action, le reste peut être effectué de manière quasi-automatique.

L’optimisation même de la vraisemblance peut être effectuée en utilisant des algorithmes divers. Cependant nous recommandons celui de Powell, étant donné qu’il n’utilise pas de gradient, et qu’il fonctionnera même en présence de sauts et autres discontinuités.

Dans une méthode alternative, on peut utiliser les mêmes filtres, mais au lieu de calculer et maximiser la vraisemblance, on augmente la dimension de l’état en y ajoutant les paramètres du modèle. Ainsi ayant par exemple quatre paramètres (comme dans le modèle de Heston) on aura un état de dimension cinq. On peut alors choisir des conditions initiales et appliquer les filtres à cet état augmenté. Après un certain nombre d’observations, les éléments de l’état correspondant aux paramètres devraient converger vers leurs valeurs réelles. Cette méthode est souvent appelée Filtre Joint (JF)
La difficulté principale de l’estimation des paramètres des modèles de volatilité stochastique, est que par sa définition même, la volatilité affecte le bruit de l’observation et non pas l’observation même. Cette remarque d’apparence anodine a des conséquences dramatiques sur la vitesse de convergence des estimateurs.

De nombreux Diagnostiques existent pour les estimateurs. Pour une série temporelle d’observations donnée, on peut après la détermination des paramètres optimaux, utiliser ceux-ci pour calculer l’erreur entre la valeur de l’observation même et la valeur prédite par le filtre. Les erreurs normalisées communément dénommées les résidus, peuvent alors être utilisées pour déterminer par exemple le variogramme. Le variogramme d’une série de points Gaussiens standards non-corrélés est constante et égale à 1.0 –sauf en zéro ou il vaut zéro par définition–

Pour les modèles de volatilité stochastique à base de diffusion, ces résidus devraient être non-corrélés et répartis selon une loi Normale standard. Des histogrammes, des tests $\chi^2$ et des tests de Box-Ljung permettent de vérifier ces propriétés.

On peut également évaluer la moyenne et l’écart type de ces résidus. Ces valeurs sont dénommées respectivement MPE et RMSE. Naturellement des valeurs élevées de MPE et RMSE correspondraient à une mauvaise performance des filtres.

Même si en réalité on ne dispose que d’un chemin historique, et par conséquent d’une série temporelle, pour juger la performance des estimateurs on devrait simuler plusieurs chemins et observer comment l’estimateur fonctionne en moyenne. Ainsi on pourra obtenir un ensemble de paramètres optimaux par chemin, et ensuite calculer la moyenne et l’écart type de ces paramètres optimaux. La distribution de ces paramètres sur les différents chemins est dénommée la distribution d’échantillonnage, et peut être obtenue via des histogrammes.

Il est clair que plus le chemin observé est long, plus les paramètres implicites obtenus seront précis. En effet les théorèmes ergodiques nous indiquent que les paramètres convergent vers leurs vraies valeurs lorsque la longueur du chemin tend vers l’infini. Cette convergence a bien lieu pour le problème de la volatilité stochastique, mais elle est encore une fois plus lente que celle des problèmes usuels.

Ceci peut être évidemment observé pour les paramètres de dérive de la volatilité, mais encore plus dramatiquement pour les paramètres de volatilité de la volatilité. En effet ces derniers n’affectent que le bruit d’un état qui lui-même n’affecte que le bruit de l’observation. Malgré ceci, en simulant des chemins suffisamment longs, on peut apercevoir que la convergence a bel et bien lieu.

Il faut cependant se souvenir qu’en réalité, on ne dispose que d’une seule série (historique) et que même si l’estimateur marchait très bien en moyenne cela ne suffirait pas pour garantir des valeurs précises. En effet les estimateurs devraient non seulement avoir un biais aussi négligeable que possible, mais ils devraient aussi être efficaces et avoir une variance aussi petite que possible.

La médiocrité relative des résultats d’inference est en accord avec les conclusions du travail effectué par des chercheurs tels que Arun Bagchi. Le fait que la volatilité
instantanée ne soit observable qu’au second ordre du développement de Taylor, est une interprétation plausible. Ceci expliquerait pourquoi ce problème n’existe pas avec les modèles de type GARCH, où la volatilité instantanée est parfaitement connue à chaque instant en fonction des prix historiques du sous-jacent. Ceci expliquerait également pourquoi choisir un intervalle temporel plus grand rendrait le processus d’inférence plus facile.

Bien que nous ayons choisi la méthodologie statistique classique, il faut au moins brièvement mentionner une autre méthode pour estimer ces paramètres, basée sur le formalisme Bayesien, où chaque paramètre est considéré non plus comme fixe mais comme une variable aléatoire. On peut donc d’abord choisir des distributions a priori pour les paramètres et par la suite les mettre à jour pour obtenir les distributions a posteriori.

Nous devrons probablement choisir les distributions a priori de ces paramètres de manière aussi générale que possible pour ne pas influencer les résultats subjectivement. Après cela, on peut par exemple considérer la moyenne a posteriori de chacun de ces paramètres pour l’estimation.

Pour certaines distributions les densités a priori et a posteriori appartiennent à la même famille. Dans ce cas, on appelle ces distributions a priori conjuguées.

Pour certains cas élémentaires, les distributions a posteriori sont analytiquement connues et par conséquent on peut calculer la moyenne et la variance des paramètres directement. Cependant en pratique, ceci n’est souvent pas possible et de ce fait on a besoin de techniques numériques de Chaîne de Markov Monte Carlo (MCMC) comme l’Échantillonneur de Gibbs où l’on simule les paramètres et l’état (la volatilité) alternativement. En outre on peut rarement simuler directement par la distribution a posteriori, et par conséquent on devra utiliser un algorithme de Metropolis-Hastings (MH) où l’on simule par une distribution candidate connue (par exemple la distribution a priori) et ensuite on accepte ou l’on rejette le point simulé en observant le quotient MH.

Ce quotient MH est simplement le quotient de la densité a posteriori et la densité candidate à l’itération \(k\), divisé par la même quantité à l’itération \(k-1\). Ce quotient (du moment qu’il est plus petit que 1.0) peut être interprété comme la probabilité avec laquelle on accepte le point simulé par la distribution candidate. On peut donc comparer cette quantité avec une variable aléatoire uniforme entre zéro et un, et si la première est plus grande que la dernière alors on accepte le point et autrement on le rejette.

Les algorithmes MCMC ont l’avantage de ne pas avoir besoin d’une optimisation de type Powell ou autre. En effet un des dangers de la méthode la maximisation de la vraisemblance est que l’optimisation pourrait nous donner un optimum local, et pour éviter cela, nous devrons choisir plusieurs conditions initiales pour nous rassurer de la convergence vers l’optimum global. Au cas où l’on obtiendrait plusieurs réponses différentes, on devra prendre la réponse qui donne la vraisemblance la plus élevée. Le
danger décrit ci-dessus devient plus important si le nombre de paramètres à estimer augmente. En effet plus la dimension du problème d’optimisation est élevée, plus la fonction de la vraisemblance devient plate.

Cependant les algorithmes MCMC sont très spécifiques au modèle que l’on choisit. Ils sont donc bien moins flexibles que les algorithmes basés sur la maximisation de la vraisemblance. En effet l’algorithme qui marche pour un modèle de Heston, est entièrement différent de celui qui fonctionnerait pour un modèle VGG. En effet la relation entre les distributions a priori et a posteriori ne sera plus la même. En revanche pour la méthode classique de maximisation de la vraisemblance via PF, cette difficulté n’existe pas.

Il est clair que lorsque l’on utilise des distributions a priori non informatives, et lorsque l’on choisit des conditions initiales quelconques, les premières valeurs simulées seront loin des valeurs réelles et c’est seulement après un certain nombre d’itérations que l’on commence à obtenir de bonnes valeurs. Cette première période que l’on ignore s’appelle la période *Burn In*. Ainsi la valeur estimée d’un paramètre serait sa moyenne en commençant après cette période. La même remarque est valable pour sa variance.

Il est intéressant de remarquer que certains utilisent une étape MH dans la méthode classique de la maximisation de vraisemblance via PF. En effet après l’étape de rééchantillonnage décrite ci-dessus, on risque d’avoir un phénomène *d’Appauvrissement d’Échantillons*, où les particules s’écroulent vers une seule valeur. Cette étape supplémentaire MH nous permet donc de régénérer les particules autour de celles qui sont importantes, i.e. celles qui ont des poids élevés et ont survécu le rééchantillonnage.

Dans tout ceci, il faut se souvenir que nous avons affaire ici à des modèles et des méthodes de calibrage paramétriques. Il existe en effet également des méthodes non-paramétriques (NP) ou encore semi-non paramétriques (SNP), où l’on utilise des formes fonctionnelles au lieu d’expressions paramétriques exactes. Aït-Sahalia et Lo d’une part et Derman et Kani d’autre, utilisent des méthodologies purement NP, où la seule hypothèse est celle de diffusion, et où la dépendance de la volatilité vis-à-vis du sous-jacent, demeure entièrement libre. D’autres comme Gallant et Tauchen utilisent dans leur *Méthode Efficiente de Moments* (EMM) des fonctions SNP.

L’avantage évident des méthodes NP/ SNP est qu’elles permettent des formulations plus générales et sont pour ainsi dire, moins contraignantes. Cela dit, même si on a plus de limitation avec les méthodes paramétriques, on a également plus de contrôle pour interpréter les résultats.
Comme évoqué précédemment, le calibrage des options nous fournit les paramètres optimaux dans un monde risque-neutre. Mais lorsque l’on utilise les séries temporelles, on obtient les paramètres dans l’univers réel.

Dans les modèles à base de diffusion, cette différence entre les mesures n’affecte que les paramètres drift de la volatilité. En effet, en utilisant le théorème de Girsanov on peut voir que la volatilité de la volatilité $\xi$ et le coefficient de la corrélation $\rho$ entre la volatilité et le rendement du prix de l’action, devraient être identiques sous les deux mesures.

Il est intéressant d’apercevoir qu’en pratique, on observe le phénomène inverse : les paramètres drift estimés sont proches sous les deux mesures, alors que les valeurs de $\xi$ et $\rho$ (en valeur absolue) sont bien plus élevées sous la mesure risque-neutre.

Ceci peut avoir deux significations : Soit les modèles de diffusion sont incorrects et l’on doit ajouter des sauts pour réconcilier les deux mesures; soit les options prédissent la possibilité d’un saut qui est absent dans la série temporelle passée.

Ce deuxième choix nous donne la possibilité d’une transaction pour profiter de cette différence. Par exemple, en achetant des options Call hors de la monnaie et en vendant des options Put hors de la monnaie, on devrait obtenir un premium qui est trop élevé si l’on considère les mouvements du sous-jacent.

Cela dit, cette transaction n’est pas sans risque. Effectivement, si jamais il y a un crash, on pourrait perdre énormément de capital. En effet dans ce cas les options auraient été correctes dans leur évaluation de risque. Ce risque perçu par certains participants mais absent dans la série historique est parfois appelé le risque de Peso.

Cette expression a été inventée par l’économiste Milton Friedman au début des années soixante-dix. Durant cette période la valeur du taux d’échange entre le dollar américain et le peso mexicain, était trop élevée et n’était pas compatible avec les niveaux relatifs des taux d’intérêts des deux pays. Cela était donc considéré comme une sorte de paradoxe à l’époque. Milton Friedman a proposé l’explication suivante : L’information contenue dans les taux d’intérêt était correcte et même si le taux d’échange était trop élevé jusqu’à ce moment, cela ne voulait pas dire qu’il resterait au même niveau éternellement. En effet peu de temps après cela, la monnaie mexicaine a été dévaluée drastiquement. Ceci a justifié a posteriori le niveau relatif des taux d’intérêt.
Le même phénomène peut en partie expliquer l’incompatibilité entre les valeurs implicites de $\xi$ contenues dans les options d’une part, et dans la série temporelle d’autre part.

La transaction décrite ci-dessus devra idéalement être faite de manière delta et vega neutre. Pour cela on devrait d’abord choisir le nombre relatif d’options Put et Call à fin d’égaliser le vega net à zéro, puis on devra choisir le nombre d’actions pour avoir le delta
total égal à zéro. Cette transaction exploite principalement la différence entre les skewness implicites dans les des densités.

On pourrait également exploiter la différence entre les kurtosis des deux distributions en effectuant la transaction suivante : On vend des options à la monnaie de même que celles loin-hors de la monnaie, puis on achète des options proche-hors de la monnaie. Notez que la première transaction utilise les différences entre les valeurs de $\xi$ et $\rho$ alors que la deuxième utilise seulement la différence entre les valeurs de $\xi$

On pourrait également utiliser les résultats de ces transactions comme une mesure empirique de la cohérence de l’information contenue dans les marchés des Options d’une part et celle du marché des Actions d’autre part. En effet si ces informations sont inconsistentes comme on le pense, alors ces transactions devraient être profitables. Si ce n’est pas le cas, cela veut dire que la différence entre les paramètres résultants de nos estimations, n’est pas fiable. Effectivement on l’a vu dans la deuxième partie, l’estimation des paramètres $\xi$ et $\rho$ via une maximisation de la vraisemblance est extrêmement difficile et par conséquent pas nécessairement crédible.

Dans le cas de modèles non-Gaussiens, comme VGG, souvent les relations d’invariance de paramètres sous les deux mesures ne seront plus valides. Cependant on peut utiliser l’information de $p(x)$ la distribution statistique obtenue des séries temporelles d’une part, et celle de la distribution risque-neutre $q(x)$ obtenue du marché des options d’autre part, de la manière suivante :

Comme l’ont démontré Carr et Madan, on peut choisir une fonction d’utilité économique et maximiser la valeur finale des investissements en se servant de cette fonction $U(x)$, le paiement final $f(x)$ à déterminer, et la distribution $p(x)$
En effet la valeur finale de la richesse sera distribuée suivant la probabilité statistique correspondant à la réalité.
Cependant ce problème d’optimisation en $f(x)$ contient des contraints sur la valeur de notre investissement initial. En effet on a initialement besoin d’au moins la valeur présente moyenne du paiement final, sous la distribution risque-neutre $q(x)$
Ainsi, on peut voir que pour déterminer la fonction de paiement optimal, on a besoin des distributions $p(x)$ et $q(x)$ à la fois, ce qui justifie leurs estimations séparées.
The Volatility Process

Introduction

Summary

This thesis focuses upon developing Methodologies for Estimating Stochastic Volatility (SV) parameters from the Stock-Price Time-Series under a Classical framework. The text contains three Chapters and is structured as follows:

In the First Chapter, we shall introduce and discuss the concept of various parametric SV models. This chapter represents a brief survey of the existing literature on the subject of non-deterministic volatility.

We start with the concept of lognormal distribution and historic volatility. We then will introduce the Black-Scholes [38] framework. We shall also mention alternative interpretations as suggested by Cox and Rubinstein [66]. We shall state how these models are unable to explain the negative-skewness and the leptokurticity commonly observed in the stock markets. Also, the famous implied-volatility smile would not exist under these assumptions.

At this point we consider the notion of level-dependent volatility as advanced by researchers such as Cox and Ross [64], [65] as well as Bensoussan, Crouhy and Galai [33]. Either an artificial expression of the instantaneous variance will be used, as is the case for Constant Elasticity Variance (CEV) models, or an implicit expression will be deduced from a Firm model similar to Merton’s [189], for instance.

We also will bring up the subject of Poisson Jumps [190] in the distributions providing a negative-skewness and larger kurtosis. These jump-diffusion models offer a link between the volatility smile and credit phenomena.

We then discuss the idea of Local Volatility [36] and its link to the instantaneous unobservable volatility. Work by researchers such as Dupire [89], Derman and Kani [74] will be cited. We shall also describe the limitations of this idea due to an ill-poised inversion phenomenon, as revealed by Avellaneda [16] and others.

Unlike Non-Parametric Local Volatility models, Parametric Stochastic Volatility (SV) models [140] define a specific stochastic differential equation for the unobservable instantaneous variance. We therefore will introduce the notion of two-factor Stochastic Volatility and its link to one-factor Generalized Autoregressive Conditionally Heteroskedastic (GARCH) processes [40]. The SV model class is the one we shall focus upon. Studies by scholars such as Engle [94], Nelson [194] and Heston [134] will be discussed at this juncture. We will briefly mention related works on Stochastic Implied Volatility by Schonbucher [213] as well as Uncertain Volatility by Avellaneda [17].
Having introduced SV, we then discuss the two-factor Partial Differential Equations (PDE) and the incompleteness of the markets when only cash and the underlying asset are used for hedging.

We then will examine Option Pricing techniques such as Inversion of the Fourier transform, Mixing Monte-Carlo as well as a few asymptotic pricing techniques, as explained for instance by Lewis [177].

At this point we shall tackle the subject of pure-jump models such as Madan’s Variance Gamma [182] or its variants VG with Stochastic Arrivals (VGSA) [48]. The latter adds to the traditional VG a way to introduce the volatility clustering (persistence) phenomenon. We will mention the distribution of the stock market as well as various option pricing techniques under these models. The inversion of the characteristic function is clearly the method of choice for option pricing in this context.

In the **Second Chapter** we will tackle the notion of Inference (or Parameter-Estimation) for Parametric SV models. We shall first briefly analyze the Cross-Sectional Inference and will then focus upon the Time-Series Inference.

We start with a concise description of cross-sectional estimation of SV parameters in a risk-neutral framework. A Least Squares Estimation (LSE) algorithm will be discussed. The Direction-Set optimization algorithm [204] will be also introduced at this point. The fact that this optimization algorithm does not use the gradient of the input-function is important, since we shall later deal with functions that contain jumps and are not necessarily differentiable everywhere.

We then discuss the parameter inference from a Time-Series of the underlying asset in the real world. We shall do this in a Classical (Non-Bayesian) [240] framework and in particular we will estimate the parameters via a Maximization of Likelihood Estimation (MLE) [127] methodology. We shall explain the idea of MLE, its link to the Kullback-Leibler [100] distance as well as the calculation of the Likelihood function for a two-factor SV model.

We will see that unlike GARCH models, SV models do not admit an analytic (integrated) likelihood function. This is why we will need to introduce the concept of Filtering [129].

The idea behind Filtering is to obtain the best possible estimation of a hidden state given all the available information up to that point. This estimation is done in an iterative manner in two stages: The first step is a Time Update where the prior distribution of the hidden state at a given point in time, is determined from all the past information via a Chapman-Kolmogorov equation. The
second step would then involve a Measurement Update where this prior distribution is used together with the conditional likelihood of the newest observation in order to compute the posterior distribution of the hidden state. The Bayes rule is used for this purpose. Once the posterior distribution is determined, it could be exploited for the optimal estimation of the hidden state.

We shall start with the Gaussian case where the first two moments characterize the entire distribution. For the Gaussian-Linear case, the optimal Kalman Filter (KF) \cite{129} is introduced. Its nonlinear extension, the Extended KF (EKF) is described next. A more suitable version of KF for strongly nonlinear cases, the Unscented KF (UKF) \cite{166} is also analyzed. In particular we will see how this filter is related to Kushner’s Nonlinear Filter (NLF) \cite{173}, \cite{174}.

EKF uses a first order Taylor approximation upon the nonlinear transition and observation functions, in order to bring us back into a simple KF framework. On the other hand, UKF uses the true nonlinear functions without any approximation. It however supposes that the Gaussianity of the distribution is preserved through these functions. UKF determines the first two moments via integrals that are computed upon a few appropriately chosen “sigma points”. NLF does the same exact thing via a Gauss-Hermite quadrature. However NLF often introduces an extra centering step, which will avoid poor performance due to an insufficient intersection between the prior distribution and the conditional likelihood.

As we shall observe, in addition to their use in the MLE approach, the Filters above could be applied to a direct estimation of the parameters via a Joint Filter (JF) \cite{133}. The JF would simply involve the estimation of the parameters together with the hidden state via a dimension augmentation. In other words, one would treat the parameters as hidden states. After choosing initial conditions and applying the filter to an observation data set, one would then disregard a number of initial points and take the average upon the remaining estimations. This initial rejected period is known as the “burn in” period.

We will test various representations or State Space Models of the Stochastic Volatility models such as Heston’s \cite{134}. The concept of Observability \cite{205} will be introduced in this context. We will see that the parameter estimation is not always accurate given a limited amount of Daily data.

Before a closer analysis of the performance of these estimation methods, we shall introduce simulation-based Particle Filters (PF) \cite{79}, \cite{122}, which can be applied to Non-Gaussian distributions. In a PF algorithm, the Importance Sampling technique is applied to the distribution. Points are simulated via a chosen proposal distribution and the resulting weights proportional to the conditional likelihood are computed. Since the variance of these weights tends to increase over time and cause the algorithm to diverge, the simulated points go through a variance reduction technique commonly referred to as Resampling \cite{14}. During
this stage, points with too small a weight are disregarded and points with large weights are reiterated. This technique could cause a Sample Impoverishment which can be corrected via a Metropolis-Hastings Accept/Reject test. Work by researchers such as Doucet [79], Smith and Gordon [122] are cited and used in this context.

Needless to say, the choice of the proposal distribution could be fundamental in the success of the PF algorithm. The most natural choice would be to take a proposal distribution equal to the prior distribution of the hidden state. Even if this makes the computations simpler, the danger would be a non-alignment between the prior and the conditional likelihood as we previously mentioned. To avoid this, other proposal distributions taking into account the observation should be considered. The Extended PF (EPF) and the Unscented PF (UPF) [229] precisely do this by adding an extra Gaussian Filtering step to the process. Other techniques such as Auxiliary PF (APF) have been developed by Pitt and Shephard [203].

Interestingly, we will see that PF brings only marginal improvement to the traditional KF’s when applied to daily data. However, for a larger time-step where the nonlinearity is stronger, the PF does help more.

At this point we also compare the Heston model to other SV models such as the “3/2” model [177] using real market data, and we will see that the latter performs better than the former. This is in line with the findings of Engle and Ishida [95]. We can therefore apply our inference tools to perform Model Identification.

Various Diagnostics [129] are used to judge the performance of the estimation tools. Mean Price Errors (MPE) and Root Mean Square Errors (RMSE) are calculated from the residual errors. The same residuals could be submitted to a Box-Ljung test, which will allow us to see whether they still contain autocorrelation. Other tests such as the Chi-Square Normality test as well as plots of Histograms and Variograms [110] are performed.

Most importantly, for the Inference process, we back-test the tools upon artificially simulated data, and we observe that although they give the correct answer asymptotically, the results remain inaccurate for a smaller amount of data points. It is reassuring to know that these observations are in agreement with work by other researchers such as Bagchi [19].

Here, we shall attempt to find an explanation for this mediocre performance. One possible interpretation comes from the fact that in the SV problem, the parameters affect the noise of the observation and not its drift. This is doubly true of volatility-of-volatility and stock-volatility correlation, which affect the noise of the noise. We should, however, note that the product of these two parameters enters in the equations at the same level as the drift of the instantaneous variance, and it is precisely this product that appears in the skewness
of the distribution.

Indeed the instantaneous volatility is observable only at the second order of a Taylor (or Ito) expansion of the logarithm of the asset-price. This also explains why one-factor GARCH models do not have this issue. In their context the instantaneous volatility is perfectly known as a function of previous data points.

The issue therefore seems to be a low Signal to Noise Ratio (SNR). We could improve our estimation by considering additional data points. Using a high frequency (several quotes a day) for the data, does help in this context. However one needs to obtain clean and reliable data first.

Also we can see why a large time-step (e.g. yearly) makes the inference process more robust by improving the observation quality. Still, using a large time-step brings up other issues such as stronger nonlinearity as well as fewer available data points, not to mention the non-applicability of the Girasnov theorem.

We shall analyze the Sampling Distributions of these parameters over many simulations and see how unbiased and efficient the estimators are. Not surprisingly, the inefficiency remains significant for a limited amount of data.

One needs to question the performance of the actual Optimization algorithm as well. It is known that the greater the number of the parameters we are dealing with, the flatter the Likelihood function and therefore the more difficult to find a global optimum.

Nevertheless, it is important to remember that the SNR and therefore the performance of the inference tool depend on the actual value of the parameters. Indeed, it is quite possible that the real parameters are such that the inference results are accurate.

We then apply our PF to a jump-diffusion model (such as the Bates [28] model) and we will see that the estimation of the jump parameters is more robust than the estimation of the diffusion parameters. This reconfirms that the estimation of parameters affecting the drift of the observation is more reliable.

We finally apply the PF to non-Gaussian models such as VGSA [48] and we will observe similar results as for the diffusion-based models. Once again the VG parameters directly affecting the observation are easier to estimate, while the arrival rate parameters affecting the noise are more difficult to recover.

Although as mentioned we use a Classical approach, we briefly discuss Bayesian methods [34] such as Markov Chain Monte Carlo’s (MCMC) [163] including the Gibbs Sampler [55] and the Metropolis-Hastings (MH) [58] algorithm. Bayesian methods consider the parameters not as fixed numbers, but random variables
having a prior distribution. One then updates these distributions from the observations similarly to what is done in the Measurement Update step of a Filter. Sometimes the prior and posterior distributions of the parameters belong to the same family and are referred to as Conjugates. The parameters are finally estimated via an averaging procedure similar to the one employed in the JF. Whether the Bayesian methods are actually better or worse than the Classical ones has been a subject of long philosophical debate [240] and remains for the reader to decide.

Other methodologies that differ from ours are the Non-Parametric (NP) and the Semi-Non-Parametric (SNP) ones. These methods are based upon Kernel-based interpolation procedures, and have the obvious advantage of being less restrictive. However, Parametric models such as the ones used by us, offer the possibility of comparing and interpreting parameters such as drift and volatility of the instantaneous variance explicitly. Researchers such as Gallant, Tauchen [109] and Aït-Sahalia [6] use NP/ SNP approaches.

Finally, in the Third Chapter, we will apply the above Parametric Inference methodologies to a few assets and will question the Consistency of information contained in the Options markets on the one hand, and in the Stock market on the other hand.

We shall see that there seems to be an excess negative-skewness and kurtosis in the former. This is in contradiction with the Gisanov theorem for a Heston model and could mean either that the model is misspecified, or that there is a profitable transaction to be made. Another explanation could come from the Peso Theory [12] (or Crash-O-Phobia [155]) where an expectation of a so-far absent crash exists in the options markets.

Adding a jump component to the distributions helps to reconcile the volatility of volatility and correlation parameters, however remains insufficient. This is in agreement with statements made by Bakshi, Cao and Chen [20].

It is important to realize that ideally, one should compare the information embedded in the options and the evolution of the underlying asset during the life of these options. Indeed ordinary put or call options are forward (and not backward) looking. However given the limited amount of available daily data through this period, we make the assumption that the dynamics of the underlying asset do not change before and during the existence of the options. We therefore use time-series that start long before the commencement of these contracts.

This assumption allows us to consider a Skewness Trade [6] where we would exploit such discrepancies by buying Out-of-the-Money (OTM) Call Options and selling OTM Put Options. We shall see that the results are not necessarily
conclusive. Indeed, even if the trade often generates profits, occasional sudden jumps cause large losses. This transaction is therefore similar to “selling insurance”.

We also apply the same idea to the VGSA model where despite the non-Gaussian features, the volatility of the arrival rate is supposed to be the same under the real and risk-neutral worlds.

Let us be clear on the fact that this chapter does not constitute a thorough empirical study of stock versus options markets. It rather presents a set of examples of application for our previously constructed inference tools. There clearly could be many other applications, such as Model Identification as discussed in the second chapter.

Yet another application of the separate estimations of the statistical and risk-neutral distributions, is the determination of optimal positions in derivatives securities as discussed by Carr and Madan [52]. Indeed the expected Utility function to be maximized needs the real world distribution, while the initial wealth constraint exploits the risk-neutral distribution. This can be seen via a self-financing portfolio argument similar to the one used by Black and Scholes [38].

Finally, we should remember that in all the above, we are assuming that the asset and options dynamics follow a known and fixed model such as Heston or VGSA. This is clearly a simplification of reality. Indeed the true markets follow an unknown and perhaps more importantly constantly changing model. The best we can do is to use the information hitherto available and hope that the future behavior of the assets is not too different from the past one. Needless to say, as time passes by and new information becomes available, we need to update our models and parameter values. This could be done within either a Bayesian or Classical framework.

In the Bibliography section, many but not all relevant articles and books are cited. Only some of them are directly referred to in the text.
Contributions and Further Research

The Contribution of the thesis is presenting a general and systematic way to calibrate any parametric SV model (diffusion-based or not) to a Time-Series under a Classical (non-Bayesian) framework. Although the concept of Filtering has been used for estimating volatility processes before [130], to our knowledge, this has always been for specific cases and was never generalized. The use of Particle Filtering allows us to do this in a flexible and simple manner. We also studied the convergence properties of our tools and showed their limitations.

Whether the results of these calibrations are consistent with the information contained in the Options markets is a fundamental question. The applications of this test are numerous among which the Skewness trade is only one example.

What else can be done? A comparative study between our approach and Bayesian ones on the one hand, and Non-Parametric ones on the other hand. Work by researchers such as Johannes, Polson and Aït-Sahalia would be extremely valuable in this context.

Data and Programs

This thesis centers on Time-Series methodologies and exploits either artificially generated inputs, or real market data. When real market data is utilized, the source is generally Bloomberg. However, most of the data could be obtained from other public sources available on the Internet.

All numeric computations are performed via routines implemented in the C programming language. Some algorithms such as the Direction-Set Optimization algorithm are taken from “Numerical Recipes in C” [204]. No statistical packages such as S-Plus or R have been used.

The Skewness transactions studied in the third chapter of this thesis, were carried out at RBC Capital Markets and remain proprietary. All Cross-Sectional Calibrators (whether implemented via Fourier-Inversion or Monte-Carlo Mixing) belong to RBC Capital Markets or Citigroup. However the Time-Series Filters and Likelihood Maximization routines, which constitute the major portion of the text, belong to the author and are available upon request.

Scope

We study Stochastic Volatility (SV) models under a Parametric framework. We therefore do not use Non-Parametric or Semi-Non-Parametric (SNP) methodologies such as the Efficient Method of Moments [109], Indirect Inference [123]
or Kernel-based algorithms [100].

We use a Classical framework where the unknown parameters are fixed. In particular, we focus upon Likelihood Maximization algorithms. We therefore are not using Bayesian approaches such as Markov-Chain Monte-Carlo’s [163] even if we do mention and discuss them.

We are principally focusing upon two-factor SV models and not one-factor GARCH [96] models. However we do compare our approach to GARCH-based algorithms.

We use Bates and VGSA models containing jumps in the stock return, but not in the volatility process itself as Matytsin [187] does for instance.

We consider only one asset at a time and do not evaluate cross-asset transactions exploiting correlations, as is done for instance by Avellaneda [15] in the context of Dispersion Trading.

We perform separate inference analysis for the Options and the Stock Markets, since we intend to examine the consistency of the information contained in them. Others [56], [165] present a joint parameter-estimation from the two markets.

We analyze the dynamics of Equity Indices as well as individual Stocks. We do not study Foreign-Exchange or Interest-Rate Dynamics. The methodologies could however be applied to these markets without great difficulty. It is important to note that a new approach based on Chaos theory, unifying Interest-Rate Dynamics with Stochastic Volatility, was suggested by Hughston [144].

In our empirical studies we use Daily time-series and not High Frequency (HF) data as some [24] do in the context of Realized Volatility. Yet as we shall state, having access to reliable HF data would be extremely valuable.

Finally, we shall not discuss the speed of our various Filters. Indeed the computational times are highly machine dependent and could vary between a few minutes (e.g. Kalman Filter applied to 5000 data points) and several days (e.g. Particle Filter applied to 5,000,000 data points via 1000 particles). Our primary concern has been to have more accurate and not necessarily faster estimation tools.
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Chapter 1

The Volatility Problem

Suppose we use the standard deviation of possible future returns on a stock as a measure of its volatility. Is it reasonable to take that volatility as a constant over time? I think not.
- Fisher Black

1.1 Introduction

It is widely accepted today that an assumption of a constant volatility fails to explain the existence of the volatility smile as well as the leptokurtic character (fat tails) of the stock distribution. The above Fisher Black quote made shortly after the famous constant-volatility Black-Scholes model was developed, proves the point.

In this Chapter, we will start by describing the concept of Brownian Motion for the Stock Price Return as well as the concept of Historic Volatility.

We will then discuss the Derivatives Market and the ideas of Hedging and Risk Neutrality. We will briefly describe the Black-Scholes Partial Derivatives Equation (PDE) in this section.

Next, we will talk about Jumps and Level Dependent Volatility models. We will first mention the Jump Diffusion process and introduce the concept of Leverage. We will then refer to two popular Level Dependent approaches: the Constant Elasticity Variance (CEV) model and the Bensoussan-Crouhy-Galai (BCG) model.
At this point, we will mention Local Volatility models developed in the recent past by Dupire and Derman-Kani and we will discuss their stability.

Following this, we will tackle the subject of Stochastic Volatility where we will mention a few popular models such as the Square-Root model and GARCH.

We will then talk about the Pricing PDE under Stochastic Volatility and the risk-neutral version of it. For this we will need to introduce the concept of Market Price of Risk.

The Generalized Fourier Transform is the subject of the following section. This technique was used by Alan Lewis extensively for solving Stochastic Volatility problems.

Next, we will discuss the Mixing Solution, both in a correlated and non-correlated case. We will mention its link to the Fundamental Transform and its usefulness for Monte-Carlo based methods.

We will then describe the Long Term Asymptotic case, where we get closed-form approximations for many popular methods such as the Square-Root model.

We will finally talk about pure-jump models such as Variance Gamma and VGSA.
1.2 The Stock Market

1.2.1 The Stock Price Process

The relationship between the stock market and the mathematical concept of Brownian Motion goes back to Bachelier [18]. A Brownian Motion corresponds to a process the increments of which are independent stationary normal random variables. Given that a Brownian Motion can take negative values, it cannot be used for the stock price. Instead, Samuelson [211] suggested to use this process to represent the return of the stock price, which will make the stock price a Geometric (or exponential) Brownian Motion.

In other words, the stock price \( S \) follows a Log-Normal process

\[
\text{d}S_t = \mu S_t \text{dt} + \sigma S_t \text{d}B_t
\]

(1.1)

where \( \text{d}B_t \) is a Brownian Motion process, \( \mu \) the instantaneous expected total return of the stock (possibly adjusted by a dividend yield) and \( \sigma \) the instantaneous standard deviation of stock price returns, called the volatility in financial markets.

Using Ito’s lemma,\(^2\) we also have

\[
\text{d} \ln(S_t) = (\mu - \frac{1}{2} \sigma^2) \text{dt} + \sigma \text{d}B_t
\]

(1.2)

The stock return \( \mu \) could easily become time dependent without changing any of our arguments. For simplicity, we will often refer to it as \( \mu \) even if we mean \( \mu_t \). This remark holds for other quantities such as \( r_t \) the interest-rate, or \( q_t \) the dividend-yield.

The equation (1.1) represents a continuous process. We can either take this as an approximation to the real discrete tick by tick stock movements, or consider it the real unobservable dynamics of the stock price, in which case the discrete prices constitute a sample from this continuous ideal process. Either way, the use of a continuous equation makes the pricing of financial instruments more analytically tractable.

The discrete equivalent of (1.2) is

\[
\ln S_{t+\Delta t} = \ln S_t + (\mu - \frac{1}{2} \sigma^2) \Delta t + \sigma \sqrt{\Delta t} B_t
\]

(1.3)

where \( (B_t) \) is a sequence of independent normal random variables with zero mean and variance of 1.

\(^1\)For an introduction to Stochastic Processes see Karatzas [167] or Oksendal [197].
\(^2\)See for example Hull [146].
1.2.2 Historic Volatility

This suggests a first simple way to estimate the volatility $\sigma$, namely the *historic volatility*. Considering $S_1, ..., S_N$ a sequence of known historic daily stock close prices, calling $R_n = \ln(S_{n+1}/S_n)$ the stock price return between two days and $\bar{R} = \frac{1}{N} \sum_{n=0}^{N-1} R_n$ the mean return, the historic volatility would be the annualized standard deviation of the returns, namely

$$\sigma_{hist} = \sqrt{\frac{252}{N-1} \sum_{n=0}^{N-1} (R_n - \bar{R})^2} \quad (1.4)$$

Because we work with annualized quantities, and we are using daily stock close prices we needed the factor 252, supposing that there are approximately 252 business days in a year.\(^3\)

Note that $N$ the number of observations can be more or less than one year, hence when talking about a historic volatility, it is important to know what time horizon we are considering? We can indeed have three month historic volatility or three year historic volatility. Needless to say taking too few prices would give an inaccurate estimation. Similarly the begin and end date of the observations matter. It is preferable to take the end date as close as possible to today, so that we include more recent observations.

An alternative was suggested by Parkinson [200] where instead of daily close prices, we use the high and the low prices of the stock on that day, and $R_n = \ln(S_{high}^n/S_{low}^n)$.

The volatility would then be

$$\sigma_{parkinson} = \sqrt{\frac{252}{N-1} \left[ \frac{1}{4 \ln(2)} \right] \sum_{n=0}^{N-1} (R_n - \bar{R})^2}$$

This second moment estimation derived by Parkinson, is based upon the fact that the range $R_n$ of the asset, follows a *Feller* distribution.

Plotting for instance the one-year rolling\(^4\) historic volatility (1.4) of S&P500 Stock Index, it is easily seen that this quantity is *not* constant over time. This observation was made as early as the 60’s by many financial mathematicians and followers of the *Chaos Theory*. We therefore need time-varying volatility models.

One natural extension of the constant volatility approach is to make $\sigma_t$ a deterministic function of time. This is equivalent to giving the volatility a term structure, by analogy with interest rates.

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\(^3\)Clearly the observation frequency does not have to be daily.

\(^4\)By *rolling* we mean that the one-year interval slides within the total observation period.
Figure 1.1: SPX Historic Rolling Volatility from 2000/01/03 to 2001/12/31. As we can see, the volatility is clearly non-constant.
1.3 The Derivatives Market

Until now we only mentioned the stock price movements independently from the derivatives market, we now are going to include the financial derivatives (specially options) prices as well. These instruments became very popular and as liquid as the stocks themselves after Black and Scholes introduced their risk-neutral pricing formula in [38].

1.3.1 The Black Scholes Approach

The Black Scholes approach makes a number of reasonable assumptions about markets being frictionless and uses the log-normal model for the stock price movements. It also supposes a constant or deterministically time dependent stock drift and volatility. Under these conditions they prove that it is possible to hedge a position in a contingent claim dynamically by taking an offsetting position in the underlying stock and hence become immune to the stock movements. This risk neutrality is possible because, as they show, we can replicate the financial derivative (for instance an option) by taking positions in Cash and the Underlying Security. This condition of possibility of replication is called Market Completeness.

In this situation everything happens as if we were replacing the stock drift \( \mu_t \) with the riskfree rate of interest \( r_t \) in (1.1) or \( r_t - q_t \) if there is a dividend-yield \( q_t \). The contingent claim \( f(S,t) \) having a payoff \( G(S_T) \) will satisfy the famous Black-Scholes equation

\[
rf = \frac{\partial f}{\partial t} + (r - q)S \frac{\partial f}{\partial S} + \frac{1}{2} \sigma^2 S^2 \frac{\partial^2 f}{\partial S^2} \quad (1.5)
\]

Indeed the Hedged Portfolio \( \Pi = f - \frac{\partial f}{\partial S} S \) is immune to the stock random movements and according to Ito’s lemma verifies

\[
d\Pi = \left( \frac{\partial f}{\partial t} + \frac{1}{2} \sigma^2 S^2 \frac{\partial^2 f}{\partial S^2} \right) dt
\]

which must also be equal to \( r\Pi dt \) or else there would be possibility of Riskless Arbitrage.\(^5\)

Note that this equation is closely related to the Feynman-Kac equation satisfied by \( F(S,t) = E_t(h(S_T)) \) for any function \( h \) under the risk-neutral measure.

\(^5\)For a detailed discussion see Hull [146].
The Volatility Process

$F(S,t)$ must be a Martingale under this measure and therefore must be driftless, which implies $dF = \sigma S \frac{\partial F}{\partial S} dB_t$ and

$$0 = \frac{\partial F}{\partial t} + (r - q)S \frac{\partial F}{\partial S} + \frac{1}{2} \sigma^2 S^2 \frac{\partial^2 F}{\partial S^2}$$

This would indeed be a different way to reach the same Black-Scholes equation, by using $f(S,t) = \exp(-rt)F(S,t)$ as was done for instance in Shreve [218].

Let us insist again on the fact that the real drift of the stock price does not appear in the above equation, which makes the volatility $\sigma_t$ the only unobservable quantity.

As we said the volatility could be a deterministic function of time without changing the above argument, in which case all we need to do is to replace $\sigma^2$ with $\frac{1}{t} \int_0^t \sigma^2 ds$ and keep everything else the same.

For Calls and Puts, where the payoffs $G(S_T)$ are respectively $MAX(0, S_T - K)$ and $MAX(0, K - S_T)$ where $K$ is the strike price and $T$ the maturity of the option, the Black Scholes Partial Derivatives Equation is solvable and gives the celebrated Black Scholes formula

\begin{equation}
\text{call}_t = S_t e^{-q(T-t)} \Phi(d_1) - K e^{-r(T-t)} \Phi(d_2) \tag{1.6}
\end{equation}

and

\begin{equation}
\text{put}_t = -S_t e^{-q(T-t)} \Phi(-d_1) + K e^{-r(T-t)} \Phi(-d_2) \tag{1.7}
\end{equation}

where $\Phi(x) = \frac{1}{\sqrt{2\pi}} \int_{-\infty}^{x} e^{-\frac{u^2}{2}} du$ is the Cumulative Standard Normal function and $d_1 = d_2 + \sigma \sqrt{T - t}$ and $d_2 = \frac{\ln(S_t/K) + (r - q - \frac{1}{2} \sigma^2)(T - t)}{\sigma \sqrt{T - t}}$.

Note that using the well known symmetry property for Normal distributions $\Phi(-x) = 1 - \Phi(x)$ in the above formulae, we could reach the Put-Call Parity relationship

\begin{equation}
\text{call}_t - \text{put}_t = S_t e^{-q(T-t)} - K e^{-r(T-t)} \tag{1.8}
\end{equation}

that we can also rearrange as

$$S_t e^{-q(T-t)} - \text{call}_t = K e^{-r(T-t)} - \text{put}_t$$

the left-hand side of the last equation is called a covered call, and is equivalent to a short position in a put combined with a bond.

\^6For an explanation see Shreve [218] or Karatzas [167].
1.3.2 The Cox Ross Rubinstein Approach

Later, Cox, Ross and Rubinstein [66] developed a simplified approach using the Binomial Law to reach the same pricing formulae. The approach commonly referred to as the Binomial Tree uses a tree of recombining spot prices where at a given time step \( n \) we have \( n + 1 \) possible \( S[n][j] \) spot prices, with \( 0 \leq j \leq n \). Calling \( p \) the upward transition probability and \( 1 - p \) the downward transition probability, \( S \) the stock price today, \( S_u = uS \) and \( S_d = dS \) upper and lower possible future spot prices, we can write the expectation equation\(^7\)

\[
E[S] = puS + (1 - p)dS = e^{r\Delta t} S
\]

which immediately gives us

\[
p = \frac{a - d}{u - d}
\]

with \( a = \exp(r\Delta t) \).

We can also write the variance equation

\[
Var[S] = pu^2S^2 + (1 - p)d^2S^2 - e^{2r\Delta t} S^2 \approx \sigma^2 S^2 \Delta t
\]

which after choosing a centering condition such as \( ud = 1 \), will provide us with \( u = \exp(\sigma\sqrt{\Delta t}) \) and \( d = \exp(-\sigma\sqrt{\Delta t}) \).

Using the above values for \( u, d \) and \( p \) we can build the Tree and using the final payoff we can calculate the option price by backward induction.\(^8\)

We can also build this tree by applying an Explicit Finite Difference scheme to the PDE (1.5) as was done in Wilmott [238].

An important advantage of the Tree method is that it can be applied to American Options (with early exercise) as well.

It is possible to deduce the implied volatility of Call and Put options by solving a reverse Black-Scholes equation: \( i.e. \) find the volatility that would equate the Black-Scholes price to the market price of the option.

This is a good way to see how derivatives markets perceive the underlying volatility. It is easy to see that if we change the maturity and strike prices of options (and keep everything else fixed) the implied volatility will not be constant. It will have a linear skew and a convex form as the strike price changes. This famous “smile” cannot be explained by simple time dependence, hence the necessity of introducing new models.\(^9\)

\(^7\)The expectation equation is written under the risk-neutral probability.

\(^8\)For an in-depth discussion on Binomial Trees see Cox [67].

\(^9\)It is interesting to note that this smile phenomenon was practically non-existent prior to the 1987 stock-market crash. Many researchers therefore believe that the markets have learnt to factor-in a crash possibility, which creates the volatility smile.
Figure 1.2: SPX Volatility Smile on Feb 12 2002 with Index=1107.5 USD, 1 month and 7 months to Maturity. The negative skewness is clearly visible. Note how the Smile becomes flatter as time to Maturity increases.
1.4 Jump Diffusion and Level Dependent Volatility

In addition to the volatility smile observable from the implied volatilities of the options, there is evidence that the assumption of a pure normal distribution (also called pure diffusion) for the stock return is not accurate. Indeed “fat tails” have been observed away from the mean of the stock return. This phenomenon is called leptokurticity and could be explained in many different ways.

1.4.1 Jump Diffusion

Some try to explain the smile and the leptokurticity by changing the underlying stock distribution from a diffusion process to a jump-diffusion process. A Jump Diffusion is not a Level Dependant Volatility process, however we are mentioning it in this section to demonstrate the importance of the leverage effect.

Merton [190] was first to actually introduce jumps in the stock distribution. Kou [172] recently used the same idea to explain both the existence of fat tails and the volatility smile.

The stock price will follow a modified stochastic process under this assumption. If we add to the Brownian Motion $dB_t$ a Poisson (Jump) process $dq$ with an intensity$^{11}$ $\lambda$, then calling $k = \mathbb{E}(Y - 1)$ with $Y - 1$ the random variable percentage change in the stock price, we will have

$$dS_t = (\mu - \lambda k)S_t dt + \sigma S_t dB_t + S_t dq$$

(1.9)

or equivalently

$$S_t = S_0 \exp[(\mu - \sigma^2/2 - \lambda k)t + \sigma B_t]Y_n$$

where $Y_0 = 1$ and $Y_n = \prod_{j=1}^n Y_j$ with $Y_j$’s independently identically distributed random variables and $n$ a Poisson random variable with a parameter $\lambda t$.

It is worth noting that for the special case where the jump corresponds to total ruin or default, we have $k = -1$ which will give us

$$dS_t = (\mu + \lambda)S_t dt + \sigma S_t dB_t + S_t dq$$

(1.10)

and

$$S_t = S_0 \exp[(\mu + \sigma^2/2)t + \sigma B_t]Y_n$$

$^{10}$See for instance Karatzas [167].

$^{11}$The intensity could be interpreted as the mean number of jumps per time unit.
Given that in this case $E(Y_n) = E(Y_n^2) = e^{-\lambda t}$ it is fairly easy to see that in the risk-neutral world
$$E(S_t) = S_0 e^{rt}$$

exactly as in the pure diffusion case, but

$$Var(S_t) = S_0^2 e^{2rt} (e^{(\sigma^2 + \lambda)t} - 1) \approx S_0^2 (\sigma^2 + \lambda)t$$

(1.11)

unlike the pure diffusion case where $Var(S_t) \approx S_0^2 \sigma^2 t$.

Proof:
Indeed
$$E(S_t) = S_0 \exp((r + \lambda)t) \exp(-\frac{\sigma^2}{2}t) E[\exp(\sigma B_t)] E(Y_n) =$$
$$S_0 \exp((r + \lambda)t) \exp(-\frac{\sigma^2}{2}t) \exp(-\lambda t) = S_0 \exp(rt)$$

and
$$E(S_t^2) = S_0^2 \exp(2(r + \lambda)t) \exp(-\sigma^2 t) E[\exp(2\sigma B_t)] E(Y_n^2) =$$
$$S_0^2 \exp(2(r + \lambda)t) \exp(-\sigma^2 t) \exp(\frac{(2\sigma)^2}{2}) \exp(-\lambda t) =$$
$$S_0^2 \exp((2r + \lambda)t) \exp(\sigma^2 t)$$

and as usual
$$Var(S_t) = E(S_t^2) - E^2(S_t)$$

(QED)

Link to Credit Spread

Note that for a zero-coupon risky bond $Z$ with no recovery, a credit spread $C$ and a face value $X$ paid at time $t$ we have
$$Z = e^{-(r+C)t}X = e^{-\lambda t}(e^{-rt}X) + (1 - e^{-\lambda t})(0)$$

consequently $\lambda = C$ and using (1.11) we can write
$$\tilde{\sigma}^2(C) = \sigma^2 + C$$

where $\sigma$ is the fixed (pure diffusion) volatility and $\tilde{\sigma}$ is the modified jump diffusion volatility. The above equation relates the volatility and leverage, a concept we will see later in level dependent models as well.

Also, we could see that everything happens as if we were using the Black-Scholes
pricing equation but with a modified “interest rate” which is $r + C$. Indeed the Hedged Portfolio $\Pi = f - \frac{\partial f}{\partial S} S$ now satisfies

$$d\Pi = \left( \frac{\partial f}{\partial t} + \frac{1}{2} \sigma^2 S^2 \frac{\partial^2 f}{\partial S^2} \right) dt$$

under the no-default case which occurs with a probability of $e^{-\lambda dt} \approx 1 - \lambda dt$ and

$$d\Pi = -\Pi$$

under the default case which occurs with a probability of $1 - e^{-\lambda dt} \approx \lambda dt$.

We therefore have

$$E(d\Pi) = \left( \frac{\partial f}{\partial t} + \frac{1}{2} \sigma^2 S^2 \frac{\partial^2 f}{\partial S^2} - \lambda \Pi \right) dt$$

and using a diversification argument we can always say that $E(d\Pi) = r \Pi dt$ which provides us with

$$(r + \lambda) f = \frac{\partial f}{\partial t} + (r + \lambda) S \frac{\partial f}{\partial S} + \frac{1}{2} \sigma^2 S^2 \frac{\partial^2 f}{\partial S^2}$$

which again is the Black-Scholes PDE with a “risky rate”.

A generalization of the Jump Diffusion process would be the use of the Lévy Process. A Lévy Process is a stochastic process with independent and stationary increments. Both the Brownian Motion and the Poisson process are included in this category. For a description, see Matacz [186].

### 1.4.2 Level Dependent Volatility

Many assume that the smile and the fat tails are due to the level dependence of the volatility.

The idea would be to make $\sigma_t$ level dependent or a function of the spot itself, we would therefore have

$$dS_t = \mu_t S_t dt + \sigma(S, t) S_t dB_t$$

(1.13)

Note that to be exact, a Level Dependent Volatility is a function of the spot price alone. When the volatility is a function of the spot price and time, it is referred to as Local Volatility, which we shall discuss further.

The Constant Elasticity Variance Approach

One of the very first attempts to use this approach was the Constant Elasticity Variance (CEV) method realized by Cox [64] and [65]. In this method we would suppose an equation of the type

$$\sigma(S, t) = CS_t^\gamma$$

(1.14)
where $C$ and $\gamma$ are parameters to be calibrated either from the stock price returns themselves or from the option prices and their implied volatilities. The CEV method was recently analyzed by Jones [165] in a paper where he uses two $\gamma$ exponents.

This level depending volatility represents an important feature that is observed in options markets as well as in the underlying prices: The negative correlation between the stock price and the volatility, also called the Leverage Effect.

The Bensoussan Crouhy Galai Approach

Bensoussan, Crouhy and Galai (BCG) [33] try to find the level dependence of the volatility differently from Cox and Ross. Indeed in the CEV model, Cox and Ross first suppose that $\sigma(S, t)$ has a certain exponential form and only then try to calibrate the model parameters to the market. BCG, on the other hand try to deduce the functional form of $\sigma(S, t)$ by using a firm structure model.

The idea of firm structure is not new and goes back to Merton [189] where he considers that the Firm Assets follow a Log-Normal process

$$dV = \mu_V V dt + \sigma_V V dB_t$$

where $\mu_V$ and $\sigma_V$ are the assets return and volatility. One important point is that $\sigma_V$ is considered constant.

Merton then argues that the equity $S$ of the firm could be considered a call option on the assets of the firm with a strike price $K$ equal to the face value of the firm liabilities and an expiration $T$ equal to the average liability maturity.

Using Ito’s lemma, it is fairly easy to see that

$$dS = \mu S dt + \sigma(S, t) S dB_t = \left( \frac{\partial S}{\partial t} + \mu_V V \frac{\partial S}{\partial V} + \frac{1}{2} \sigma_V^2 V^2 \frac{\partial^2 S}{\partial V^2} \right) dt + \sigma_V V \frac{\partial S}{\partial V} dB_t$$

which immediately provides us with

$$\sigma(S, t) = \sigma_V \frac{V \partial S}{S \partial V}$$

which is an implicit functional form for $\sigma(S, t)$.

BCG then eliminate the asset term in the above functional form and end up with a non-linear PDE

$$\frac{\partial \sigma}{\partial t} + \frac{1}{2} \sigma^2 S^2 \frac{\partial^2 \sigma}{\partial S^2} + (r + \sigma^2) S \frac{\partial \sigma}{\partial S} = 0$$

This PDE gives the dependence of $\sigma$ on $S$ and $t$.

Proof:
A quick sketch of the proof is as follows: $S$ being a contingent-claim on $V$ we have the risk-neutral Black-Scholes PDE

$$\frac{\partial S}{\partial t} + rV \frac{\partial S}{\partial V} + \frac{1}{2} \sigma^2 V^2 \frac{\partial^2 S}{\partial V^2} = rS$$

and using $\frac{\partial S}{\partial V} = 1/\frac{\partial V}{\partial S}$ as well as $\frac{\partial S}{\partial t} = -\frac{\partial S}{\partial V} \frac{\partial V}{\partial t}$ and $\frac{\partial^2 S}{\partial V^2} = -\frac{\partial^2 V}{\partial S^2}/\left(\frac{\partial V}{\partial S}\right)^3$ we have the reciprocal Black-Scholes equation

$$\frac{\partial V}{\partial t} + rS \frac{\partial V}{\partial S} + \frac{1}{2} \sigma^2 S^2 \frac{\partial^2 V}{\partial S^2} = rV$$

Now posing $\Psi(S, t) = \ln V(S, t)$ we have $\frac{\partial V}{\partial t} = V \frac{\partial \Psi}{\partial t}$ as well as $\frac{\partial V}{\partial S} = V \frac{\partial \Psi}{\partial S}$ and $\frac{\partial^2 V}{\partial S^2} = V \left(\frac{\partial^2 \Psi}{\partial S^2} + \left(\frac{\partial \Psi}{\partial S}\right)^2\right)$ we will have the new PDE

$$r = \frac{\partial \Psi}{\partial t} + rS \frac{\partial \Psi}{\partial S} + \frac{1}{2} \sigma^2 S^2 \left(\frac{\partial^2 \Psi}{\partial S^2} + \left(\frac{\partial \Psi}{\partial S}\right)^2\right)$$

and the equation

$$\sigma = \sigma V/(S \frac{\partial \Psi}{\partial S})$$

this last identity implies $\frac{\partial \Psi}{\partial S} = \frac{\partial S}{\partial \Psi}$ as well as $\frac{\partial^2 \Psi}{\partial S^2} = -\frac{\sigma V(\sigma + S \frac{\partial \sigma}{\partial S})}{S \sigma^2}$ and therefore the PDE becomes

$$r = \frac{\partial \Psi}{\partial t} + r\sigma V/\sigma + \frac{1}{2} \left(\sigma^2 V - \sigma \left(\sigma + S \frac{\partial \sigma}{\partial S}\right)\right)$$

taking the derivative with respect to $S$ and using $\frac{\partial^2 \Psi}{\partial S^2} = -\frac{\sigma V \frac{\partial \sigma}{\partial S}}{S \sigma^2}$ we get the final PDE

$$\frac{\partial \sigma}{\partial t} + \frac{1}{2} \sigma^2 S^2 \frac{\partial^2 \sigma}{\partial S^2} + (r + \sigma^2) S \frac{\partial \sigma}{\partial S} = 0$$

as previously stated. \(QED\)

We therefore have an implicit functional form for $\sigma(S, t)$ and just like for the CEV case, we need to calibrate the parameters to the market data.
Figure 1.3: CEV Model for SPX on Feb 12 2002 with Index=1107.5 USD, 1 month to Maturity. The smile is fitted well, but the model assumes a perfect (negative) correlation between the stock and the volatility.
Figure 1.4: BCG Model for SPX on Feb 12 2002 with Index=1107.5 USD, 1 month to Maturity. The smile is fitted well.
1.5 Local Volatility

In the early 90’s Dupire [89] on the one hand, and Derman & Kani [74] on the other, developed a concept called Local Volatility where the volatility smile was retrieved from the option prices.

1.5.1 The Dupire Approach

The Breeden & Litzenberger Identity

This approach, uses the options prices to get the implied distribution for the underlying stock. To do this we can write

\[ V(S_0, K, T) = \text{call}(S_0, K, T) = e^{-rT} \int_0^{+\infty} (S - K)^+ p(S_0, S, T) dS \] (1.19)

where \( S_0 \) is the stock price at time \( t = 0 \) and \( K \) the strike price of the call, and \( p(S_0, S, T) \) the unknown transition density for the stock-price. As usual \( x^+ = \text{MAX}(x, 0) \)

Using the above equation and differentiating with respect to \( K \) twice we get the Breeden & Litzenberger [44] implied distribution

\[ p(S_0, K, T) = e^{rT} \frac{\partial^2 V}{\partial K^2} \] (1.20)

Proof:

The proof is straightforward if we write

\[ e^{rT} V(S_0, K, T) = \int_K^{+\infty} Sp(S_0, S, T) dS - K \int_K^{+\infty} p(S_0, S, T) dS \]

and take the first derivative

\[ e^{rT} \frac{\partial V}{\partial K} = -Kp(S_0, K, T) + Kp(S_0, K, T) - \int_K^{+\infty} p(S_0, S, T) dS \]

and the second derivative in the same manner. (QED)

The Dupire Identity

Now according to the Fokker-Planck (or forward Kolmogorov) equation\(^\text{12}\) for this density, we have

\[ \frac{\partial p}{\partial T} = \frac{1}{2} \frac{\partial^2 (\sigma^2(S,t)S^2p)}{\partial S^2} - r \frac{\partial (Sp)}{\partial S} \]

\(^\text{12}\)See for example Wilmott [237] for an explanation on Fokker-Planck equation.
and therefore after a little rearrangement
\[ \frac{\partial V}{\partial T} = \frac{1}{2} \sigma^2 K^2 \frac{\partial^2 V}{\partial K^2} - rK \frac{\partial V}{\partial K} \]
which provides us with the local volatility formula
\[ \sigma^2(K, T) = \frac{\frac{\partial V}{\partial T} + rK \frac{\partial V}{\partial K}}{\frac{1}{2} K^2 \frac{\partial^2 V}{\partial K^2}} \]  
(1.21)

Proof:
For a quick proof of the above let us use the zero interest rates case (the general case could be done similarly). we would then have
\[ p(S_0, K, T) = \frac{\partial^2 V}{\partial K^2} \]
as well as Fokker-Planck
\[ \frac{\partial p}{\partial T} = \frac{1}{2} \frac{\partial^2 \left( \sigma^2(S, t)S^2p \right)}{\partial S^2} \]
Now
\[ \frac{\partial V}{\partial T} = \int_0^{+\infty} (S_T - K)^+ \frac{\partial p}{\partial T} dS_T = \int_0^{+\infty} (S_T - K)^+ \frac{1}{2} \frac{\partial^2 \left( \sigma^2(S, T)S^2p \right)}{\partial S^2} dS_T \]
and integrating by parts twice and using the fact that
\[ \frac{\partial^2 (S_T - K)^+}{\partial K^2} = \delta(S_T - K) \]
with \( \delta(.) \) the Dirac function, we will have
\[ \frac{\partial V}{\partial T} = \frac{1}{2} \sigma^2(K, T)K^2 p(S_0, K, T) = \frac{1}{2} K^2 \sigma^2(K, T) \frac{\partial^2 V}{\partial K^2} \]
as stated. (QED)

It is also possible to use the implied volatility \( \sigma_{BS} \) from the Black-Scholes formula (1.6) and express the above local volatility in terms of \( \sigma_{BS} \) instead of \( V \). For a detailed discussion we could refer to Wilmott [237].

Local Volatility vs. Instantaneous Volatility

Clearly the local volatility is related to the instantaneous variance \( \nu \), as Gatheral [113] shows, the relationship could be written as
\[ \sigma^2(K, T) = \mathbb{E}[\nu_T | S_T = K] \]  
(1.22)
that is, local variance is the risk-neutral expectation of the instantaneous variance conditional on the final stock price being equal to the strike price.\footnote{13}{Note that this is independent from the process for \( \nu_t \), meaning that any stochastic volatility model satisfies this property, which is an attractive feature of local volatility models.}

**Proof:** Let us show the above identity for the case of zero interest rates.\footnote{14}{For the case of non-zero rates we need to work with the Forward Price instead of the Stock Price.}

As mentioned above, we have

\[
\sigma^2(K, T) = \frac{\partial V}{\partial T} \frac{1}{2} K^2 \frac{\partial^2 V}{\partial K^2}
\]

On the other hand using the call payoff \( V(S_0, K, t = T) = \mathbb{E}[(S_T - K)^+] \) we have

\[
\frac{\partial V}{\partial K} = \mathbb{E}[H(S_T - K)]
\]

with \( H(.) \) the Heaviside function and

\[
\frac{\partial^2 V}{\partial K^2} = \mathbb{E}[\delta(S_T - K)]
\]

with \( \delta(.) \) the Dirac function.

Therefore the Ito lemma at \( t = T \) would provide

\[
d(S_T - K)^+ = H(S_T - K) dS_T + \frac{1}{2} \nu_T S_T^2 \delta(S_T - K) dT
\]

Using the fact that the Forward Price (here with zero interest rates, the Stock Price) is a Martingale under the risk-neutral measure

\[
dV = d\mathbb{E}[(S_T - K)^+] = \frac{1}{2} \mathbb{E}[\nu_T S_T^2 \delta(S_T - K)] dT
\]

Now we have

\[
\mathbb{E}[\nu_T S_T^2 \delta(S_T - K)] = \mathbb{E}[\nu_T | S_T = K] K^2 \mathbb{E}[\delta(S_T - K)] = \mathbb{E}[\nu_T | S_T = K] K^2 \frac{\partial^2 V}{\partial K^2}
\]

Putting all this together

\[
\frac{\partial V}{\partial T} = \frac{1}{2} K^2 \frac{\partial^2 V}{\partial K^2} \mathbb{E}[\nu_T | S_T = K]
\]

and by the above expression of \( \sigma^2(K, T) \) we will have

\[
\sigma^2(K, T) = \mathbb{E}[\nu_T | S_T = K]
\]

as claimed. (QED)
1.5.2 The Derman Kani Approach

The Derman & Kani technique is very similar to the above approach, except it uses the Binomial (or Trinomial) Tree framework instead of the continuous one. Using the Binomial Tree notations, their upward transition probability \( p_i \) from the spot \( s_i \) at time \( t_n \) to the upper node \( S_{i+1} \) at the following time-step \( t_{n+1} \), is obtained from the usual

\[
p_i = \frac{F_i - S_i}{S_{i+1} - S_i}
\]

where \( F_i \) is the stock forward price known from the market, and \( S_i \) the lower spot at the step \( t_{n+1} \).

In addition, we have for a call expiring at time-step \( t_{n+1} \)

\[
C(K, t_{n+1}) = e^{-r\Delta t} \sum_{j=1}^{n} [\lambda_j p_j + \lambda_{j+1} (1 - p_{j+1})] \text{MAX}(S_{j+1} - K, 0)
\]

where \( \lambda_j \)’s are the known Arrow-Debreu prices corresponding to the discounted probability of getting to the point \( s_j \) at time \( t_n \) from \( S_0 \) the initial stock price. These probabilities could easily be derived iteratively. This allows us after some calculation to obtain \( S_{i+1} \) as a function of \( s_i \) and \( S_i \), namely

\[
S_{i+1} = S_i [e^{r\Delta t} C(s_i, K, t_{n+1}) - \Sigma] - \lambda_i (F_i - S_i)
\]

where the term \( \Sigma \) represents the sum \( \sum_{j=i+1}^{n} \lambda_j (F_j - s_j) \). This means that after choosing the usual centering condition for the Binomial Tree

\[
s_i^2 = S_i S_{i+1}
\]

we have all the elements to build the tree and deduce the implied distribution from the Arrow-Debreu prices.

1.5.3 Stability Issues

The local volatility models are very elegant and theoretically sound, however they present in practice many stability issues. They are Ill-Posed Inversion problems and are extremely sensitive to the input data.\(^{15}\) This might introduce arbitrage opportunities and in some cases negative probabilities or variances. Derman and Kani suggest overwriting techniques to avoid such problems. Andersen [13] tries to improve this issue by using an Implicit Finite Difference method, however he recognizes that the negative variance problem could still happen.

\(^{15}\)See Tavella [226] or Avellaneda [16].
One way to make the results smoother is to use a constrained optimization. In other words, when trying to fit theoretical results $C_{\text{theo}}$ to the market prices $C_{\text{mrkt}}$ instead of minimizing

$$\sum_{j=1}^{N} (C_{\text{theo}}(K_j) - C_{\text{mrkt}}(K_j))^2$$

we could minimize

$$\lambda \frac{\partial \sigma}{\partial t} + \sum_{j=1}^{N} (C_{\text{theo}}(K_j) - C_{\text{mrkt}}(K_j))^2$$

where $\lambda$ is a constraint parameter which could also be interpreted as a Lagrange multiplier. However this is an artificial way to smoothen the results and the real issue remains that once again, we have an inversion problem that is inherently unstable.

What is more, local volatility models imply that future implied volatility smiles will be flat relative to today’s, which is another limitation. As we will see in the following section, Stochastic Volatility models offer more time-homogeneous volatility smiles.

An alternative approach suggested in [16] would be to choose a prior risk-neutral distribution for the asset (based on a subjective view) and then minimize the relative Entropy distance between the desired surface and this prior distribution. This approach uses the Kullback-Leibler distance (which we will discuss in the context of MLE) and performs the minimization via Dynamic Programming [35] on a tree.

### 1.5.4 Calibration Frequency

One of the most attractive features of Local-Vol models is their ability to match plain-vanilla puts and calls exactly. This will avoid arbitrage situations, or worse, market manipulations by traders to create “phantom” profits.

As explained in Hull [147], these arbitrage-free models were developed by researchers with a Single Calibration (SC) methodology assumption. However traders use them with a Continual Recalibration (CR) strategy in practice. Indeed if they used the SC version of the model, significant errors would be introduced from one week to the following as shown by Dumas et al. [88].

However once this CR version is used, there is no guarantee that the no-arbitrage solution

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16See Gatheral [114].
property of the original SC model is preserved. Indeed the Dupire equation determines the marginal stock distribution at different points in time, but not the joint distribution of these stock prices. Therefore a path-dependent option could very well be mispriced, and the more path-dependent this option, the greater the mispricing.

Hull [147] takes the example of a Bet Option, a Compound Option and a Barrier Option. The Bet Option depends on the distribution of the stock at one point in time and therefore is correctly priced with a Continually Recalibrated Local-Vol model. The Compound Option has some path dependency, hence a certain amount of mispricing compared to a Stochastic Volatility (SV) model. Finally, the Barrier Option has a strong degree of path dependency and will introduce large errors.

Note that this is due to the discrete nature of the data. Indeed the maturities we have are limited. If we had all possible maturities in a continuous way, the joint distribution would be determined completely.

Also, when interpolating in time, it is customary to interpolate upon the true variance $t \sigma_t^2$ rather than the volatility $\sigma_t$ given the equation

$$T_2\sigma^2(T_2) = T_1\sigma^2(T_1) + (T_2 - T_1)\sigma^2(T_1, T_2)$$

Interpolating upon the true variance will provide smoother results as shown by Jackel [152].

Proof:

Indeed, calling for $0 \leq T_1 \leq T_2$ the spot return variances

$$Var(0, T_2) = T_2\sigma^2(T_2)$$

$$Var(0, T_1) = T_1\sigma^2(T_1)$$

for a Brownian Motion, we have independent increments and therefore a forward variance $Var(T_1, T_2)$ such that

$$Var(0, T_1) + Var(T_1, T_2) = Var(0, T_2)$$

which demonstrates the point. (QED)
1.6 Stochastic Volatility

Unlike Non-Parametric Local Volatility models, Parametric Stochastic Volatility (SV) models define a specific stochastic differential equation for the unobservable instantaneous variance. As we shall see, the previously defined CEV model could be considered a special case of these models.

1.6.1 Stochastic Volatility Processes

The idea would be to use a different stochastic process for $\sigma$ altogether. Making the volatility a deterministic function of the spot is a special “degenerate” two-factor, a natural generalization of which would precisely be to have two stochastic processes with a non-perfect correlation.\(^\text{17}\)

Several different stochastic processes have been suggested for the volatility. One popular one is the Ornstein-Uhlenbeck (OU) process:

$$d\sigma_t = -\alpha \sigma_t dt + \beta dZ_t$$  \hspace{1cm} (1.24)

where $\alpha$ and $\beta$ are two parameters, remembering the stock equation

$$dS_t = \mu_t S_t dt + \sigma_t S_t dB_t$$

there is a (usually negative) correlation $\rho$ between $dZ_t$ and $dB_t$ which can in turn be time or level dependent.

Heston [134] and Stein [223] were among those who suggested the use of this process. Using Ito’s lemma, we can see that the stock-return variance $v_t = \sigma_t^2$ satisfies a Square-Root or Cox-Ingersoll-Ross (CIR) process

$$dv_t = (\omega - \theta v_t) dt + \xi \sqrt{v_t} dZ_t$$  \hspace{1cm} (1.25)

with $\omega = \beta^2$, $\theta = 2\alpha$ and $\xi = 2\beta$.

Note that the OU process has a closed form solution

$$\sigma_t = \sigma_0 e^{-\alpha t} + \beta \int_0^t e^{-\alpha(t-s)}dZ_s$$

\(^{17}\)Note that here, the instantaneous volatility is stochastic. Recent work by researchers such as Schonbucher supposes a Stochastic Implied-Volatility process, which is a completely different approach. See for instance [213].

On the other hand, Avellaneda et al. [17] use the concept of Uncertain Volatility for pricing and hedging Derivative Securities. They make the volatility switch between two extreme values based on the convexity of the derivative contract and obtain a nonlinear Black-Scholes-Barenblatt equation, which they solve on a grid.
which means that $\sigma_t$ follows in law $\Phi(\sigma_0 e^{-\alpha t}, \frac{\beta^2}{2\alpha}(1 - e^{-2\alpha t}))$, with $\Phi$ again the Normal Distribution. This was discussed in Fouque [104] and Shreve [218].

Heston and Nandi [137] show that this process corresponds to a special case of the General Auto-Regressive Conditional Heteroskedasticity (GARCH) model that we will discuss further below.

Another popular process is the GARCH(1,1) process where we would have

$$dv_t = (\omega - \theta v_t)dt + \xi v_t dZ_t$$ (1.26)

### 1.6.2 GARCH and Diffusion Limits

The most elementary GARCH process called GARCH(1,1) was developed originally in the field of econometrics by Engle [94] and Bollerslev [40] in a discrete framework. The stock discrete equation (1.3) could be rewritten by taking $\Delta t = 1$ and $v_n = \sigma_n^2$ as

$$\ln S_{n+1} = \ln S_n + (\mu - \frac{1}{2} v_{n+1}) + \sqrt{v_{n+1}} B_{n+1}$$ (1.27)

calling the mean adjusted return

$$u_n = \ln \left( \frac{S_n}{S_{n-1}} \right) - (\mu - \frac{1}{2} v_n) = \sqrt{v_n} B_n$$ (1.28)

calling the variance process in GARCH(1,1) is supposed to be

$$v_{n+1} = \omega_0 + \beta v_n + \alpha u_n^2 = \omega_0 + \beta v_n + \alpha v_n B_n^2$$ (1.29)

where $\alpha$ and $\beta$ are weight parameters and $\omega_0$ a parameter related to the long-term variance.\(^{18}\)

Nelson [194] shows that as the time interval length decreases and becomes infinitesimal, the above equation (1.29) becomes precisely the previously cited equation (1.26). To be more accurate there is a weak convergence of the discrete GARCH process to the continuous diffusion limit.\(^{19}\) For a GARCH(1,1) continuous diffusion, the correlation between $dZ_t$ and $dB_t$ is zero.

It might appear surprising that even if the GARCH(1,1) process has only one

\(^{18}\)It is worth mentioning that as explained in [100] a GARCH(1,1) model could be rewritten as an Auto-Regressive Moving Average model of first order ARMA(1,1) and therefore an Auto-Regressive model of infinite order AR(+\(\infty\)). GARCH is therefore a parsimonious model that can fit the data with only a few parameters. Fitting the same data with an ARCH or AR model would require a much larger number of parameters. This feature makes the GARCH model very attractive.

\(^{19}\)For an explanation on weak convergence see for example Varadhan [230].
source of randomness, namely $B_n$, the continuous version has two independent Brownian Motions. This is understandable if we consider $B_n$ a standard normal random variable and $A_n = B_n^2 - 1$ another random variable. It is fairly easy to see that $A_n$ and $B_n$ are uncorrelated even if $A_n$ is a function of $B_n$. As we go towards the continuous version, we can use Donsker’s Theorem,20 by letting the time interval $\Delta t \to 0$, to prove that we end up with two uncorrelated and therefore independent Brownian Motions. This is a limitation of the GARCH(1,1) model, hence the introduction of the Non-Linear Asymmetric GARCH (NGARCH) model.

Duan [83] attempts to explain the volatility smile by using the NGARCH process expressed by

$$v_{n+1} = \omega_0 + \beta v_n + \alpha (u_n - c\sqrt{v_n})^2$$

(1.30)

where $c$ is a parameter to be determined.

The NGARCH process was first introduced by Engle [97]. The continuous counterpart of NGARCH is the same equation (1.26) except unlike the equation resulting from GARCH(1,1) there is a non-zero correlation between the stock process and the volatility process. This correlation is created precisely because of the parameter $c$ that was introduced, and is once again called the leverage effect. The parameter $c$ is sometimes referred to as the leverage parameter.

We can find the following relationships between the discrete process and the continuous diffusion limit parameters by letting the time interval become infinitesimal

$$\omega = \frac{\omega_0}{dt^2}$$

$$\theta = \frac{1 - \alpha(1 + c^2) - \beta}{dt}$$

$$\xi = \alpha \sqrt{\frac{\kappa - 1 + 4c^2}{dt}}$$

and the correlation between $dB_t$ and $dZ_t$

$$\rho = \frac{-2c}{\sqrt{\kappa - 1 + 4c^2}}$$

where $\kappa$ represents the Pearson Kurtosis21 of the mean adjusted returns ($u_n$). As we can see, the sign of the correlation $\rho$ is determined by the parameter $c$.

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20For a discussion on Donsker’s Theorem, similar to the Central Limit Theorem, see for instance Whitt [235].
21The Kurtosis corresponds to the Fourth Moment. The Pearson Kurtosis for a Normal Distribution is equal to 3.
Proof:
A quick proof of the convergence to diffusion limit could be outlined as follows. Let us assume that \( c = 0 \) for simplicity, we therefore are dealing with the GARCH(1,1) case. As we saw
\[
v_{n+1} = \omega_0 + \beta v_n + \alpha v_n B_n^2
\]
then
\[
v_{n+1} - v_n = \omega_0 + \beta v_n - v_n + \alpha v_n - \alpha v_n + \alpha v_n B_n^2
\]
or
\[
v_{n+1} - v_n = \omega_0 - (1 - \alpha - \beta)v_n + \alpha v_n (B_n^2 - 1)
\]
Now, allowing the time-step \( \Delta t \) to become variable and posing
\[
Z_n = \frac{(B_n^2 - 1)}{\sqrt{\kappa - 1}} v_n + \Delta t - v_n = \omega \Delta t^2 - \theta \Delta t v_n + \xi v_n \sqrt{\Delta t} Z_n
\]
and annualizing \( v_n \) by posing \( v_t = v_n/\Delta t \) we shall have
\[
v_{t+\Delta t} - v_t = \omega \Delta t - \theta \Delta t v_t + \xi v_t \sqrt{\Delta t} Z_n
\]
and as \( \Delta t \to 0 \) we get
\[
dv_t = (\omega - \theta v_t) dt + \xi v_t dZ_t
\]
as claimed. (QED)

Note that the discrete GARCH version of the Square-Root process (1.25) is
\[
v_{n+1} = \omega_0 + \beta v_n + \alpha (B_n - c \sqrt{v_n})^2
\]
(1.31)
as Heston and Nandi show\(^{22}\) in [137].

Also, note that having a diffusion process \( dv_t = b(v_t) dt + a(v_t) dZ_t \) we can apply an Euler Approximation\(^{23}\) to discretize and obtain a Monte-Carlo process such as
\[
v_{n+1} - v_n = b(v_n) \Delta t + a(v_n) \sqrt{\Delta t} Z_n
\]
It is important to note that if we use a GARCH process and go to the continuous diffusion limit, and then apply an Euler Approximation, we will not necessarily find the original GARCH process again. Indeed, there are many different ways to discretize the continuous diffusion limit and the GARCH process corresponds to one special way.

In particular, if we use (1.31) and allow \( \Delta t \to 0 \) to get to the continuous diffusion limit, we shall obtain (1.25). As we will see later in the section on Mixing Solutions, we can then apply a discretization to this process and obtain a Monte-Carlo simulation
\[
v_{n+1} = v_n + (\omega - \theta v_n) \Delta t + \xi \sqrt{v_n} \sqrt{\Delta t} Z_n
\]
\(^{22}\) For a detailed discussion on the convergence of different GARCH models towards their Diffusion Limits, also see Duan [85].
\(^{23}\) See for instance Jones [165].
which is again different from (1.31) but obviously has to be consistent in terms of pricing. However, we should know which method we are working with from the very beginning to perform our calibration on the corresponding specific process.

Corradi [61] explains this in the following manner: The discrete GARCH model could converge either towards a two-factor continuous limit if one chooses the Nelson parameterization, or could very well converge to a one-factor diffusion limit if one chooses another parameterization. What is more, an appropriate Euler discretization of the one-factor continuous model will provide a GARCH discrete process, while as previously mentioned the discretization of the two-factor diffusion model provides a two-factor discrete process. This distinction is fundamental and could explain why GARCH and SV behave differently in terms of parameter estimation.
Figure 1.5: GARCH Monte-Carlo Simulation with the Square-Root Model for SPX on Feb 12 2002 with Index=1107.5 USD, 1 month to Maturity. Powell Optimization method was used for Least-Square Calibration.
1.7 The Pricing PDE under Stochastic Volatility

A very important issue to underline here is that, because of the unhedgeable second source of randomness, the concept of Market Completeness is lost. We can no longer have a straightforward risk neutral pricing. This is where the market price of risk will come into consideration.

1.7.1 The Market Price of Volatility Risk

Indeed, taking a more general form for the variance process

\[ dv_t = b(v_t)dt + a(v_t)dZ_t \]  

(1.32)

as we previously said, using the Black-Scholes risk-neutrality argument, the equation (1.1) could be replaced with

\[ dS_t = (r_t - q_t)S_tdt + \sigma_tS_tdB_t \]  

(1.33)

This is equivalent to changing the probability measure from the real one to the risk neutral one.

We therefore need to use (1.33) together with the risk adjusted volatility process

\[ dv_t = \tilde{b}(v_t)dt + a(v_t)dZ_t \]  

(1.34)

where

\[ \tilde{b}(v_t) = b(v_t) - \lambda a(v_t) \]

with \( \lambda \) the market price of volatility risk. This quantity is closely related to the market price of risk for the stock \( \lambda_e = (\mu - r)/\sigma \). Indeed as Hobson [140] and Lewis [177] both show, we have

\[ \lambda = \rho \lambda_e + \sqrt{1 - \rho^2} \lambda^* \]  

(1.35)

where \( \lambda^* \) is the market price of risk associated with \( dB_t - \rho dZ_t \) which can also be regarded as the market price of risk for the hedged portfolio.

The passage from the above equation (1.32) to (1.34) and the introduction of the market price of volatility risk could also be explained by the Girsanov theorem as was done for instance in Fouque [104].

It is important to underline the difference between the real and the risk neutral measures here. If we use historic stock prices together with the real stock-return drift \( \mu \) to estimate the process parameters, we will obtain the real volatility-drift

\[^{24}\text{See Hull [146] or Shreve [218] for more detail.}\]
An alternative method would be to estimate \( \tilde{b}(v) \) by using current option prices and performing a Least Square Estimation. These Calibration methods will be discussed in more detail in the following chapters.

The risk neutral version for a discrete NGARCH model would also involve the market price of risk and instead of the usual

\[
\ln S_{n+1} = \ln S_n + \left( \mu - \frac{1}{2} v_{n+1} \right) + \sqrt{v_{n+1}} B_{n+1}
\]

we would have

\[
\ln S_{n+1} = \ln S_n + \left( r - \frac{1}{2} v_{n+1} \right) + \sqrt{v_{n+1}} \tilde{B}_{n+1}
\]

(1.36)

\[
v_{n+1} = \omega_0 + \beta v_n + \alpha v_n (B_n - c)^2
\]

where \( \tilde{B}_n = B_n + \lambda_e \) which could be regarded as the discrete version of the Girsanov theorem. Note that the market price of risk for the stock \( \lambda_e \) is not separable from the leverage parameter \( c \) in the above formulation.

Duan shows in [84] and [86] that the above risk-neutral GARCH system (1.36) will indeed converge towards the continuous risk-neutral GARCH

\[
dS_t = S_t r dt + S_t \sqrt{v_t} dB_t
\]

\[
dv_t = (\omega - \tilde{\theta} v_t) dt + \xi v_t dZ_t
\]

as we expected.

1.7.2 The Two Factor PDE

From here, writing a two factor PDE for a derivative security \( f \) becomes a simple application of the two-dimensional Itô’s lemma. The PDE will be\(^{25}\)

\[
r f = \frac{\partial f}{\partial t} + (r - q) S \frac{\partial f}{\partial S} + \frac{1}{2} v S^2 \frac{\partial^2 f}{\partial S^2} + \tilde{b}(v) \frac{\partial f}{\partial v} + \frac{1}{2} a^2(v) \frac{\partial^2 f}{\partial v^2} + \rho a(v) \sqrt{v} \frac{\partial^2 f}{\partial S \partial v}
\]

(1.37)

Therefore it is possible, after calibration, to apply a Finite Difference Method\(^{26}\) to the above PDE to price the derivative \( f(S, t, v) \). An alternative would be to use directly the stochastic processes for \( dS_t \) and \( dv_t \) and apply a Two-Factor Monte-Carlo simulation. Later in the chapter we will also mention other possible methods such as the Mixing Solution or Asymptotic Approximations.

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\(^{25}\)For a proof of the derivation see Wilmott [237] or Lewis [177].

\(^{26}\)See for instance Tavella [227] or Wilmott [237] for a discussion on Finite Difference Methods.
Other possible approaches for incomplete markets and stochastic volatility assumption include Super-Replication and Local Risk Minimization.\textsuperscript{27} The Super-Replication strategy is the cheapest self-financing strategy with a terminal value no less than the payoff of the Derivative contract. This technique was primarily developed by El-Karoui and Quenez in [91]. Local Risk Minimization involves a partial hedging of the risk. The risk is reduced to an “intrinsic component” by taking an offsetting position in the underlying security as usual. This method was developed by Follmer and Sondermann in [102].

\footnote{For a discussion on both these techniques see Frey [107].}
1.8 The Generalized Fourier Transform

1.8.1 The Transform Technique

One useful technique to apply to the PDE (1.37) is the Generalized Fourier Transform. First, we can use the variable \( x = \ln S \) in which case, using Ito’s lemma, (1.37) could be rewritten as

\[
rf = \frac{\partial f}{\partial t} + (r - q - \frac{1}{2}v) \frac{\partial f}{\partial x} + \frac{1}{2}v \frac{\partial^2 f}{\partial x^2} + \tilde{b}(v) \frac{\partial f}{\partial v} + \frac{1}{2}a^2(v) \frac{\partial^2 f}{\partial v^2} + \rho a(v) \sqrt{v} \frac{\partial^2 f}{\partial x \partial v}
\]  

(1.38)

Calling

\[
\hat{f}(k, v, t) = \int_{-\infty}^{+\infty} e^{ikx} f(x, v, t) dx
\]

(1.39)

where \( k \) is a complex number, \( \hat{f} \) will be defined in a complex strip where the imaginary part of \( k \) is between two real numbers \( \alpha \) and \( \beta \). Once \( \hat{f} \) is suitably defined, meaning that \( k_i = \mathcal{I}(k) \) (the imaginary part of \( k \)) is within the appropriate strip, we can write the Inverse Fourier Transform

\[
f(x, v, t) = \frac{1}{2\pi} \int_{ik_1}^{ik_1 + \infty} e^{-ikx} \hat{f}(k, v, t) dk
\]

(1.40)

where we are integrating for a fixed \( k \) parallel to the real axis.

Each derivative satisfying (1.37) or equivalently (1.38) has a known payoff \( G(S_T) \) at maturity. For instance, as we said before, a Call Option has a payoff \( \text{MAX}(0, S_T - K) \) where \( K \) is the call strike-price. It is easy to see that for \( k_i > 1 \) the Fourier Transform of a call option exists and the payoff Transform is

\[
\frac{-K^{ik+1}}{k^2 - ik}
\]

(1.41)

\textbf{Proof:}

Indeed, we can write

\[
\int_{-\infty}^{+\infty} e^{ikx} (e^x - K)^+ dx = \int_{\ln K}^{+\infty} e^{ikx} (e^x - K) dx = 0 - \left( \frac{K^{ik+1}}{ik+1} - K^{ik} \right)
\]

\[28\text{See Lewis [177] for a detailed discussion on this technique.}

\[29\text{As usual we note } i = \sqrt{-1}.\]
\[-K^{ik+1}\left(\frac{1}{ik+1} - \frac{1}{ik}\right) = -K^{ik+1}\frac{1}{k^2 - ik}\]

as stated. \((QED)\)

The same could be applied to a Put option or other derivative securities. In particular, a Covered Call (Stock minus Call) having a payoff \(\min(S_T, K)\) will have a transform for \(0 < k_i < 1\) equal to

\[
\frac{K^{ik+1}}{k^2 - ik}
\]

(1.42)

Applying the Transform to the PDE (1.38) and introducing \(\tau = T - t\) and

\[
\hat{h}(k, v, \tau) = e^{(r+ik(r-q))\tau} \bar{f}(k, v, \tau)
\]

(1.43)

and posing

\[
c(k) = \frac{1}{2}(k^2 - ik)
\]

we get the new PDE equation

\[
\frac{\partial \hat{h}}{\partial \tau} = \frac{1}{2} v^2(a(v)) \frac{\partial^2 \hat{h}}{\partial v^2} + (\hat{b}(v) - ik \rho(a(v)) \sqrt{v}) \frac{\partial \hat{h}}{\partial v} - c(k)v\hat{h}
\]

(1.44)

Lewis calls the **Fundamental Transform** a function \(\hat{H}(k, v, \tau)\) satisfying the PDE (1.44) and satisfying the initial condition \(\hat{H}(k, v, \tau = 0) = 1\).

If we know this Fundamental Transform, we can then multiply it by the derivative security’s payoff transform and then divide it by \(e^{(r+ik(r-q))\tau}\) and apply the inverse Fourier technique by keeping \(k_i\) in an appropriate strip and finally get the derivative as a function of \(x = \ln S\).

### 1.8.2 Special Cases

There are cases where the Fundamental Transform is known. The case of a constant (or deterministic) volatility is the most elementary one. Indeed using (1.44) together with \(dv_t = 0\) we can easily find

\[
\hat{H}(k, v, \tau) = e^{-c(k)v\tau}
\]

which is analytic in \(k\) over the entire complex plane. Using the Call payoff Transform (1.41) we can rederive the Black-Scholes equation. The same can be done if we have a deterministic volatility \(dv_t = b(v)dt\) by using the function \(Y(v, t)\) where \(dY = b(Y)dt\).

The Square-Root model (1.25) is another important case where \(\hat{H}(k, v, \tau)\) is known and analytic. We have for this process

\[
dv_t = (\omega - \theta v_t)dt + \xi \sqrt{v_t}dZ_t
\]

30We are following Lewis [177] notations.
or under the risk-neutral measure

\[ dv_t = (\omega - \tilde{\theta}v_t)dt + \xi\sqrt{v_t}dZ_t \]

with \( \tilde{\theta} = (1 - \gamma)\rho \xi + \sqrt{\theta^2 - \gamma(1 - \gamma)\xi^2} \) where \( \gamma \leq 1 \) represents the risk-aversion factor.

For the Fundamental Transform, we get

\[ \hat{H}(k, v, \tau) = \exp[f_1(t) + f_2(t)v] \] (1.45)

with

\[ t = \frac{1}{2}\xi^2\tau \quad \tilde{\omega} = \frac{2}{\xi}\omega \quad \tilde{c} = \frac{2}{\xi}c(k) \quad \text{and} \]

\[ f_1(t) = [tg - \ln(1 - h\epsilon t\tau)]\tilde{\omega} \]

\[ f_2(t) = \left[ \frac{1 - e^{\tau d}}{1 - he^{\tau d}} \right]g \]

where

\[ d = \sqrt{\theta^2 + 4\tilde{c}} \quad g = \frac{1}{2}(\tilde{\theta} + d) \quad h = \frac{\tilde{\theta} + d}{\theta - d} \quad \text{and} \]

\[ \tilde{\theta} = \frac{2}{\xi^2}[(1 - \gamma + ik)\rho \xi + \sqrt{\theta^2 - \gamma(1 - \gamma)\xi^2}] \]

The above transform has a cumbersome expression, but it can be seen that it is analytic in \( k \) and therefore always exists. For a proof of the above refer to Lewis [177].

The Inversion of the Fourier Transform for the Square-Root (Heston) model is a very popular and powerful approach. It is appealing by its robustness and speed.

The following example is based upon SPX options as of 20040309 expiring in 1 to 8 years from the calibration date.

As we shall see further, the optimal Heston parameter-set to fit this surface could be found via a Least Square Estimation approach and for the index at \( S = 1156.86 \) USD we find the optimal parameters \( \hat{\psi}_0 = 0.1940 \) and

\[ \hat{\Psi} = (\hat{\omega}, \hat{\theta}, \hat{\xi}, \hat{\rho}) = (0.052042332, 1.8408, 0.4710, -0.4677) \]
The Volatility Process

<table>
<thead>
<tr>
<th>T / M</th>
<th>0.70</th>
<th>0.80</th>
<th>0.85</th>
<th>0.90</th>
<th>0.95</th>
<th>1.00</th>
<th>1.05</th>
<th>1.10</th>
<th>1.15</th>
<th>1.20</th>
<th>1.30</th>
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<tr>
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<td>20.16</td>
<td>18.69</td>
<td>17.96</td>
<td>17.28</td>
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<td>15.97</td>
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<td>17.29</td>
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<td>16.04</td>
<td>15.82</td>
<td>15.40</td>
</tr>
</tbody>
</table>

Table 1.1: SPX implied surface as of 03/09/2004. \( T \) is the maturity and \( M = K/S \) the inverse of the moneyness.

<table>
<thead>
<tr>
<th>T / M</th>
<th>0.70</th>
<th>0.80</th>
<th>0.85</th>
<th>0.90</th>
<th>0.95</th>
<th>1.00</th>
<th>1.05</th>
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<td>11.35</td>
<td>9.70</td>
<td>6.97</td>
</tr>
</tbody>
</table>

Table 1.2: Heston prices fitted to the 2004/03/09 surface.
Figure 1.6: SPX implied surface as of 03/09/2004. We can observe the negative skewness as well as the flattening of the slope with maturity.
1.9 The Mixing Solution

1.9.1 The Romano Touzi Approach

The idea of *Mixing Solutions* was probably presented for the first time by Hull and White [149] for a zero correlation case. Later, Romano and Touzi [209] generalized this approach for a correlated case.

The basic idea is to *separate* the random processes of the stock and the volatility, integrate the stock process conditionally upon a given volatility and finally end up with a one-factor problem.

Let us remind the two processes we had:

\[ dS_t = (r_t - q_t)S_t dt + \sigma_t S_t dB_t \]

and

\[ dv_t = \tilde{b}(v_t) dt + a(v_t) dZ_t \]

under a risk-neutral measure.

Given a correlation \( \rho_t \) between \( dB_t \) and \( dZ_t \) we can introduce the Brownian Motion \( dW_t \) independent of \( dZ_t \) and write the usual Cholesky\(^{31}\) factorization:

\[ dB_t = \rho_t dZ_t + \sqrt{1 - \rho_t^2} dW_t \]

We can then introduce the same \( X_t = \ln S_t \) and write the new system of equations:

\[ dX_t = (r - q) dt + dY_t - \frac{1}{2}(1 - \rho_t^2) \sigma_t^2 dt + \sqrt{1 - \rho_t^2} \sigma_t dW_t \quad (1.46) \]

\[ dY_t = -\frac{1}{2} \rho_t^2 \sigma_t^2 dt + \rho_t \sigma_t dZ_t \]

\[ dv_t = \tilde{b} dt + a dZ_t \]

where once again, the two Brownian Motions above are independent.

It is now possible to integrate the Stock process for a given volatility and end up with an expectation on the volatility process only. We can think of (1.46) as the limit of a discrete process, while the time step \( \Delta t \to 0 \).

For a derivative security \( f(S_0, v_0, T) \) with a payoff\(^{32}\) \( G(S_T) \), using the bivariate normal density for two uncorrelated variables, we can write

\[ f(S_0, v_0, T) = e^{-rT} E_0[ G(S_T) ] = \quad (1.47) \]

\(^{31}\)See for example Press [204].

\(^{32}\)The payoff should *not* depend on the volatility process.
\[e^{-rT} \lim_{\Delta t \to 0} \int_{-\infty}^{\infty} \ldots \int_{-\infty}^{\infty} G(S_T) \prod_{t=0}^{T-\Delta t} \exp\left[ -\frac{1}{2} (Z_t^2 + W_t^2) \right] dZ_t dW_t \]

Now if we know how to integrate the above over \(dW_t\) for a given volatility and we know the result \(f^*(S,v,T)\) (for instance for a European Call Option, we know the Black-Scholes formula (1.6), there are many other cases where we have closed-form solutions) then we can introduce the auxiliary variables

\[S^{eff} = S_0 e^{Y_T} = S_0 \exp\left( -\frac{1}{2} \int_0^T \rho_t^2 \sigma_t^2 dt + \int_0^T \rho_t \sigma_t dZ_t \right) \quad (1.48)\]

and

\[\nu^{eff} = \frac{1}{T} \int_0^T (1 - \rho_t^2) \sigma_t^2 dt \quad (1.49)\]

and as Romano and Touzi prove in [209] we will have

\[f(S_0, v_0, T) = E_0[f^*(S^{eff}, \nu^{eff}, T)] \quad (1.50)\]

where this last expectation is being taken on \(dZ_t\) only.

Note that in the zero correlation case discussed by Hull and White [149] we have \(S^{eff} = S_0\) and \(\nu^{eff} = \nu_T = \frac{1}{T} \int_0^T \sigma_t^2 dt\) which makes the expression (1.50) a natural weighted average.

### 1.9.2 A One Factor Monte-Carlo Technique

As Lewis suggests this will enable us to run a single-factor Monte-Carlo simulation on the \(dZ_t\) and apply the known closed-form for each simulated path. The method does suppose however that the payoff \(G(S_T)\) does not depend on the volatility.

Indeed, going back to (1.46) we can do a simulation on \(Y_t\) and \(v_t\) using the random sequence of \((Z_t)\), then after one path is generated, we can calculate \(S^{eff} = S_0 \exp(Y_T)\) and \(\nu^{eff} = \frac{1}{T} \sum_{t=0}^{T-\Delta t} (1 - \rho_t^2) v_t \Delta t\) and then apply the known closed-form (e.g. Black-Scholes for a call or put) with \(S^{eff}\) and \(\nu^{eff}\). Repeating this procedure for a large number of times and averaging over the paths, as we usually do in Monte-Carlo methods, we will have \(f(S_0, v_0, T)\).

This will give us a way to calibrate the model parameters to the market data. For instance using the Square-Root model

\[dv_t = (\omega - \theta v_t) dt + \xi \sqrt{v_t} dZ_t\]

we can estimate \(\omega, \theta, \xi\) and \(\rho\) from the market prices via a least-square estimation applied to theoretical prices obtained from the above Monte-Carlo

\[\text{33 Again, all notations are taken from Lewis [177].}\]
method. We can either use a single calibration and suppose we have time-independent parameters, or perform one calibration per maturity. The single calibration method is known to provide a bad fit, hence the idea of adding jumps to the stochastic volatility process as described by Matytsin [187]. However this method will introduce new parameters for calibration.\footnote{Eraker \textit{et al.} [98] claim that a model containing jumps in the return \textit{and} the volatility process will fit the options and the underlying data well, and will have no misspecification left.}
Figure 1.7: Mixing Monte-Carlo Simulation with the Square-Root Model for SPX on Feb 12 2002 with Index=1107.5 USD, 1 month and 7 months to Maturity Powell Optimization method was used for Least-Square Calibration. As we can see, both maturities are fitted fairly well.
1.10 The Long Term Asymptotic Case

In this section we will discuss the case where the contract time to maturity is very large, \( t \to \infty \). We will focus on the special case of a Square-Root process, since this is the model we will use in many cases.

1.10.1 The Deterministic Case

We shall start with the case of Deterministic Volatility and use that for the more general case of the Stochastic Volatility.

We know that under the Square-Root model the variance follows

\[
dv_t = (\omega - \theta v_t)dt + \xi \sqrt{v_t}dZ_t
\]

As an approximation we can drop the stochastic term and obtain

\[
\frac{dv_t}{dt} = \omega - \theta v_t
\]

which is an ordinary differential equation providing us immediately with

\[
v_t = \frac{\omega}{\theta} + \left(v - \frac{\omega}{\theta}\right)e^{-\theta t}
\]

where \( v \) is the initial variance for \( t = 0 \).

Using the results from the Fundamental Transform for a Covered Call Option and Put-Call parity we have for \( 0 < k_1 < 1 \)

\[
call(S, v, \tau) = Se^{-r\tau} - Ke^{-r\tau} \frac{1}{2\pi} \int_{ik_1 - \infty}^{ik_1 + \infty} e^{-ikX} \hat{H}(k, v, \tau) \frac{k}{k^2 - ik} dk
\]

where \( \tau = T - t \) and \( X = \ln(\frac{Se^{-r\tau}}{Ke^{-r\tau}}) \) represents the adjusted moneyness of the option. For the special “At The Money”\(^{35}\) case where \( X = 0 \) we have

\[
call(S, v, \tau) = Ke^{-r\tau} \left[1 - \frac{1}{2\pi} \int_{ik_1 - \infty}^{ik_1 + \infty} \frac{\hat{H}(k, v, \tau)}{k^2 - ik} dk\right]
\]

As we previously said for a deterministic volatility case, we know the Fundamental Transform

\[
\hat{H}(k, v, \tau) = \exp[-c(k)U(v, \tau)]
\]

\(^{35}\)This is different from the usual definition of At The Money calls where \( S = K \). This vocabulary is borrowed from Alan Lewis.
with \( U(v, \tau) = \int_0^\tau v(t) dt \) and as before \( c(k) = \frac{1}{2}(k^2 - i k) \), which in the special case of the Square-Root model (1.51), will provide us with

\[
U(v, \tau) = \frac{\omega}{\theta} \tau + (v - \frac{\omega}{\theta})(1 - e^{-\theta \tau})
\]

This shows once again that \( \hat{H}(k) \) is analytic in \( k \) over the entire complex plane.

Now if we let \( \tau \to \infty \) we can write the approximation

\[
\frac{\text{call}(S, v, \tau)}{K e^{-\tau r}} \approx 1 - \frac{1}{2\pi} \int_{i k_0 - \infty}^{i k_0 + \infty} \exp[-c(k)\frac{\omega}{\theta} \tau - c(k)\frac{1}{\theta} (v - \frac{\omega}{\theta})] \frac{dk}{k^2 - i k} \tag{1.54}
\]

We can either calculate the above integral exactly using the Black-Scholes theory, or take the minimum where \( c'(k_0) = 0 \), meaning \( k_0 = \frac{i}{2} \), and perform a Taylor approximation parallel to the real axis around the point \( k = k_r + \frac{i}{2} \) which will give us

\[
\frac{\text{call}(S, v, \tau)}{K e^{-\tau r}} \approx 1 - \frac{2}{\pi} \exp(-\frac{\omega}{8\theta} \tau) \exp[-\frac{1}{8\theta} (v - \frac{\omega}{\theta})] \int_{-\infty}^\infty \exp(-k^2 \frac{\omega}{2\theta} \tau) dk \]

the integral being a Gaussian we will get the result

\[
\frac{\text{call}(S, v, \tau)}{K e^{-\tau r}} \approx 1 - \sqrt{\frac{8\theta}{\pi \omega \tau}} \exp[-\frac{1}{8\theta} (v - \frac{\omega}{\theta})] \exp(-\frac{\omega}{8\theta} \tau) \tag{1.55}
\]

which finishes our deterministic approximation case.

### 1.10.2 The Stochastic Case

For the Stochastic Volatility case, Lewis uses the same Taylor expansion. He notices that for the deterministic case we had

\[
\hat{H}(k, v, \tau) = \exp[-c(k)U(v, \tau)] \approx \exp[-\lambda(k)\tau]u(k, v)
\]

for \( \tau \to \infty \) where

\[
\lambda(k) = c(k)\frac{\omega}{\theta}
\]

and

\[
u(k, v) = \exp[-c(k)\frac{1}{\theta} (v - \frac{\omega}{\theta})]
\]

If we suppose that the above identity holds for the Stochastic Volatility case as well, we can use the PDE (1.44) and interpret the result as an eigenvalue-eigenfunction identity with the eigenvalue \( \lambda(k) \) and the eigenfunction \( u(k, v) \).
This assumption is reasonable since the first Taylor approximation term for the stochastic process is deterministic.

Indeed introducing the operator
\[
\Lambda(u) = -\frac{1}{2} a^2(v) \frac{d^2 u}{dv^2} - \left[ \hat{b}(v) - ik\rho(a(v)\sqrt{v}) \right] \frac{du}{dv} + c(k)u
\]
we have
\[
\Lambda(u) = \lambda(k)u
\] (1.56)

Now the idea would be to perform a Taylor expansion around the minimum \( k_0 \) where \( \lambda'(k_0) = 0 \). Lewis shows that such \( k_0 \) is always situated on the imaginary axis. This property is referred to as the “ridge” property.

The Taylor expansion along the real axis could be written as
\[
\lambda(k) = \lambda(k_0 + k_\tau) \approx \lambda(k_0) + \frac{1}{2} k_\tau^2 \lambda''(k_0)
\]

Note that we are dealing with a minimum and therefore \( \lambda''(k_0) > 0 \). Using the above second order approximation for \( \lambda(k) \) we get
\[
\frac{\text{call}(S, v, \tau)}{K e^{-r\tau}} \approx 1 - \frac{u(k_0, v)}{k_0^2 - ik_0} \frac{1}{\sqrt{2\pi\lambda''(k_0)}} \exp[-\lambda(k_0)\tau]
\]

We can then move from the special “At The Money” case to the general case by reintroducing \( X = \ln\left( \frac{S - q\tau}{Ke^{-r\tau}} \right) \) and we will finally obtain
\[
\frac{\text{call}(S, v, \tau)}{K e^{-r\tau}} \approx e^X - \frac{u(k_0, v)}{k_0^2 - ik_0} \frac{1}{\sqrt{2\pi\lambda''(k_0)}} \exp[-\lambda(k_0)\tau - ik_0 X] \quad (1.57)
\]
which completes our determination of the asymptotic closed-form in the general case.

For the special case of the Square-Root model, taking the risk-neutral case \( \gamma = 1 \), we have\(^{36}\)
\[
\lambda(k) = -\omega g^*(k) = \frac{\omega}{\xi^2} [\sqrt{(\theta + ik\rho\xi)^2 + (k^2 - ik)\xi^2} - (\theta + ik\rho\xi)]
\]
which also allows us to calculate \( \lambda''(k) \). Also
\[
u(k, v) = \exp[g^*(k)v]
\]
where we use the notations from (1.45) and we pose
\[
g^* = g - d
\]
\(^{36}\)We can go back to the general case \( \gamma \leq 1 \) by replacing \( \theta \) with \( \sqrt{\theta^2 - \gamma(1-\gamma)\xi^2} + (1-\gamma)\rho\xi \) since this transformation is independent from \( k \) altogether.
The $k_0$ such that $\lambda'(k_0) = 0$ is

$$k_0 = \frac{i}{1 - \rho^2} \left( \frac{1}{2} - \frac{\rho}{\xi} \left( \theta - \frac{1}{2} \sqrt{4\theta^2 + \xi^2 - 4\rho\theta\xi} \right) \right)$$

which together with (1.57) provides us with the result for $\text{call}(S,v,\tau)$ in the asymptotic case under the Square-Root stochastic volatility model. Note that for $\xi \to 0$ and $\rho \to 0$ we find again the deterministic result $k_0 \to \frac{i}{2}$. 
1.10.3 A Series Expansion on Volatility-of-Volatility

Another asymptotic approach for the stochastic volatility model suggested by Lewis [177] is a Taylor expansion on the volatility-of-volatility. There are two possibilities for this, we can perform the expansion either for the Option price or for the implied-volatility directly.

In what follows we consider the former approach. Once again, we use the fundamental transform $H(k, V, \tau) = H(k, V, 0) = 1$ and

$$\frac{\partial H}{\partial \tau} = \frac{1}{2} a^2(v) \frac{\partial^2 H}{\partial v^2} + (\bar{b}(v) - ik\rho(v)\sqrt{v}) \frac{\partial H}{\partial v} - c(k)vH$$

and $c(k) = \frac{1}{2}(k^2 - ik)$.

We then pose $a(v) = \xi\eta(v)$ and expand $H(k, V, \tau)$ on powers of $\xi$ and finally apply the inverse Fourier Transform to obtain an expansion on the Call Price. With our usual notations $\tau = T - t$, $X = \ln(S/K) + (r - q)\tau$ and $Z(V) = V\tau$, the series will be

$$C(S, V, \tau) = c_{BS}(S, v, \tau) + \xi \tau^{-1}J_1\tilde{R}_{11} \frac{\partial c_{BS}(S, v, \tau)}{\partial V} +$$

$$\xi^2[\tau^{-2}J_2\tilde{R}_{20} + \tau^{-1}J_4\tilde{R}_{12} + \frac{1}{2}\tau^{-2}J_2^2\tilde{R}_{22}] \frac{\partial c_{BS}(S, v, \tau)}{\partial V} + O(\xi^3)$$

where $v(V, \tau)$ is the deterministic variance

$$v(V, \tau) = \frac{\omega}{\theta} + (V - \frac{\omega}{\theta})\left(\frac{1 - e^{-\theta\tau}}{\theta\tau}\right)$$

and $\tilde{R}_{pq} = R_{pq}(X, v(V, \tau), \tau)$ with $R_{pq}$ given polynomials of $(X, Z)$ of degree four at most, and $J_n$'s known functions of $(V, \tau)$.

The explicit expressions for all these functions are given in the third chapter of the Lewis book [177].

The obvious advantage of this approach is its speed and stability. The issue of lack of time-homogeneity of the parameters $\Psi = (\omega, \theta, \xi, \rho)$ could be addressed by performing one calibration per time-interval. In this case, for each time-interval $[t_n, t_{n+1}]$ we will have one set of parameters $\Psi_n = (\omega_n, \theta_n, \xi_n, \rho_n)$ and depending on what maturity $T$ we are dealing with, we will use one or the other parameter-set.

We compare the values obtained from this series-based approach with those from a mixing Monte-Carlo method in figure 1.8. We are taking the example that Heston studied in [134]. The graph shows the difference $C(S, V, \tau) - c_{BS}(S, V, \tau)$ for a fixed $K = 100$ USD and $\tau = 0.50$ years. The other inputs are $\omega = 0.02$, $\theta = 2.00$, $\xi = 0.10$, $\rho = -0.50$, $V = 0.01$ and $r = q = 0$. 
As we can see, the true value of the Call is lower than the Black-Scholes value for the OTM region. The higher \( \xi \) and \(|\rho|\) are, the larger this difference will be.

In figures 1.9 and 1.10 we reset the correlation \( \rho \) to zero to have a symmetric distribution, but we use a volatility-of-volatility of \( \xi = 0.10 \) and \( \xi = 0.20 \) respectively. As discussed, the parameter \( \xi \) is the one creating the leptokurticity phenomenon. A higher volatility-of-volatility causes higher valuation for Far-from-the-Money options.\(^{37}\)

\(^{37}\)Also note that the gap between the closed-form series and the Monte-Carlo model increases with \( \xi \). Indeed the accuracy of the expansion decreases as \( \xi \) becomes larger.
Figure 1.8: Comparing the Volatility-of-Volatility Series expansion with the Monte-Carlo Mixing Model. The graph shows the price difference $C(S, V, \tau) - c_{BS}(S, V, \tau)$. We are taking $\xi = 0.10$ and $\rho = -0.50$. This example was used in the original Heston paper.
Figure 1.9: Comparing the Volatility-of-Volatility Series expansion with the Monte-Carlo Mixing Model. The graph shows the price difference $C(S, V, \tau) - c_{BS}(S, V, \tau)$. We are taking $\xi = 0.10$ and $\rho = 0$. This example was used in the original Heston paper.
Figure 1.10: Comparing the Volatility-of-Volatility Series expansion with the Monte-Carlo Mixing Model. The graph shows the price difference \( C(S, V, \tau) - c_{BS}(S, V, \tau) \). We are taking \( \xi = 0.20 \) and \( \rho = 0 \). This example was used in the original Heston paper.

Unfortunately the above series approximation becomes poor as soon as the volatility of volatility becomes larger than 0.40 and the maturity becomes of the order of 1 year. This case is not unusual at all and therefore makes the use of this method limited. This is why the method of choice remains the Inversion of the Fourier Transform as previously described.
1.11 Pure-Jump Models

1.11.1 Variance Gamma

An alternative point of view is to drop the diffusion assumption altogether and replace it with a pure-jump process. Note that this is different from the Jump-Diffusion process previously discussed. Madan et al. suggested the following framework called Variance-Gamma (VG) in [182].

We would have the Log-Normal-like Stock process

\[ d\ln S_t = (\mu_S + \omega)dt + X(dt; \sigma, \nu, \theta) \]

where as before \( \mu_S \) is the real-world statistical drift of the stock log-return and \( \omega = \frac{1}{\nu} \ln(1 - \theta \nu - \sigma^2 \nu/2) \).

As for \( X(dt; \sigma, \nu, \theta) \) it has the following meaning:

\[ X(dt; \sigma, \nu, \theta) = B(\gamma(dt, 1, \nu); \theta, \sigma) \]

where \( B(dt; \theta, \sigma) \) would be a Brownian Motion with drift \( \theta \) and volatility \( \sigma \). In other words

\[ B(dt; \theta, \sigma) = \theta dt + \sigma \sqrt{dt} N(0, 1) \]

and \( N(0, 1) \) is a standard Gaussian realization.

The time-interval at which the Brownian Motion is considered is not \( dt \) but \( \gamma(dt, 1, \nu) \) which is a random realization following a Gamma distribution with a mean 1 and variance-rate \( \nu \).

The corresponding probability density function is

\[ f_{\nu}(dt, \tau) = \frac{\tau^{\frac{\nu}{2}} e^{-\frac{\tau}{\nu}}}{\nu^{\frac{\nu}{2}} \Gamma\left(\frac{\nu}{2}\right)} \]

where \( \Gamma(x) \) is the usual Gamma function.

Note that the Stock log-return density could actually be integrated for the VG model and the density of \( \ln(S_t/S_0) \) is known and could be implemented via \( K_0(x) \) the modified Bessel function of the second kind.

Indeed, calling \( z = \ln(S_k/S_{k-1}) \) and \( h = t_k - t_{k-1} \) and posing \( x_h = z - \mu_S h - \frac{h}{\nu} \ln(1 - \theta \nu - \sigma^2 \nu/2) \) we have

\[ p(z|h) = \frac{2 \exp(\theta x_h/\sigma^2)}{\nu^{1/2} \sqrt{2\pi\sigma} \Gamma\left(\frac{\nu}{2}\right)} \left( \frac{2}{2\sigma^2/\nu + \theta^2} \right)^{\frac{\nu}{4} - \frac{1}{2}} K_{\frac{\nu}{2} - \frac{1}{2}} \left( \frac{1}{\sigma^2} \sqrt{x_h^2 (2\sigma^2/\nu + \theta^2)} \right) \]

What is more, as Madan et al. show, the option valuation under VG is fairly straightforward and admits an analytically tractable closed-form which can be implemented via the above modified Bessel function of second kind and a degenerate hypergeometric function. All details are available in [182].
Remark on the Gamma Distribution

The Gamma Cumulative Distribution Function (CDF) could be defined as

$$ P(a, x) = \frac{1}{\Gamma(a)} \int_0^x e^{-t} t^{a-1} dt $$

Note that with our notations

$$ F_\nu(h, x) = F(h, x, \mu = 1, \nu) $$

with

$$ F(h, x, \mu, \nu) = \frac{1}{\Gamma\left(\frac{\mu^2 h}{\nu} \right)} \left(\frac{\mu^2 h}{\nu} \right)^\mu \int_0^x e^{-\nu \frac{\mu^2 h}{\nu} t^{\frac{\mu^2 h}{\nu} - 1} dt} $$

In other words

$$ F(h, x, \mu, \nu) = P\left(\frac{\mu^2 h}{\nu} \frac{\mu x}{\nu}\right) $$

The behavior if this CDF is displayed in figure 1.11 for different values of the parameter $a > 0$ and for $0 < x < +\infty$.

Using the inverse of this CDF we can have a simulated data-set for the Gamma law:

$$ x^{(i)} = F^{-1}_\nu(h, U^{(i)}[0, 1]) $$

with $1 \leq i \leq N_{\text{sims}}$ and $U^{(i)}[0, 1]$ a uniform random realization between zero and one.

Stochastic Volatility vs. Time-Changed processes

As mentioned in [23], this alternative formulation leading to Time-Changed processes, is closely related to the previously discussed Stochastic Volatility approach in the following way:

Taking the above VG Stochastic Differential Equation

$$ d\ln S_t = (\mu_S + \omega) dt + \theta \gamma(dt, 1, \nu) + \sigma \sqrt{\gamma(dt, 1, \nu)} N(0, 1) $$

one could consider $\sigma^2 \gamma(t, 1, \nu)$ as the integrated-variance and define $v_t(\nu)$ the instantaneous-variance as

$$ \sigma^2 \gamma(dt, 1, \nu) = v_t(\nu) dt $$

in which case, we would have

$$ d\ln S_t = (\mu_S + \omega) dt + \theta/\sigma^2 v_t(\nu) dt + \sqrt{v_t(\nu)} dt N(0, 1) = $$

$$ (\mu_S + \omega + (\theta/\sigma^2) v_t(\nu)) dt + \sqrt{v_t(\nu)} dZ_t $$

where $dZ_t$ is a Brownian Motion.

This last equation is a traditional Stochastic Volatility one.
Figure 1.11: The Gamma Cumulative Distribution Function $P(a, x)$ for various values of the parameter $a$. The implementation is based on code available in “Numerical Recipes in C”.
The Modified Bessel Function of Second Kind

Figure 1.12: The Modified Bessel Function of Second kind for a given parameter. The implementation is based on code available in “Numerical Recipes in C”.
Figure 1.13: The Modified Bessel Function of Second kind as a function of the parameter. The implementation is based on code available in “Numerical Recipes in C”.
1.11.2 Variance Gamma with Stochastic Arrival

An extension to the VG model would be a Variance Gamma model with Stochastic Arrival (VGSA) which would include the volatility Clustering effect. This phenomenon (also represented by GARCH) means that a high (low) volatility will be followed by a series of high (low) volatilities.

In this approach we replace the $dt$ in the previously defined $f_{\nu}(dt, \tau)$ with $y_t dt$ where $y_t$ follows a Square-Root (CIR) process

$$dy_t = \kappa(\eta - y_t)dt + \lambda \sqrt{y_t} dW_t$$

where the Brownian Motion $dW_t$ is independent from other processes in the model.

This is therefore a VG process where the arrival time itself is stochastic. The mean-reversion of the Square-Root process will cause the volatility persistence effect that is empirically observed. Note that (not counting $\mu_S$) the new model parameter-set is $\Psi = (\kappa, \eta, \nu, \theta, \sigma)$.

Option Pricing under VGSA

The option pricing could be carried out via a Monte-Carlo simulation algorithm under the risk-neutral measure, where as before $\mu_S$ is replaced with $r - q$. We first would simulate the path of $y_t$ by writing

$$y_k = y_{k-1} + \kappa(\eta - y_{k-1})\Delta t + \lambda \sqrt{y_{k-1}} \sqrt{\Delta t} Z_k$$

then calculate

$$Y_T = \sum_{k=0}^{N-1} y_k \Delta t$$

and finally apply one-step simulations

$$T^* = F^{-1}_\nu(Y_T, U[0,1])$$

and\footnote{This means that $T$ in VG, is replaced with $Y_T$. The rest remains identical.}

$$\ln S_T = \ln S_0 + (r - q + \omega)T + \theta T^* + \sigma \sqrt{T^*} B_k$$

Note that we have two Normal random-variables $B_k$, $Z_k$ as well as a Gamma-distributed random variable $T^*$, and that they are all uncorrelated.

Once the Stock price $S_T$ is properly simulated, we can calculate the option price as usual.
The Characteristic Function

As previously discussed, another way to tackle the option-pricing issue would be to use the characteristic functions.

For VG, the characteristic function is

$$\Psi(u, t) = \mathbb{E}[e^{iuX(t)}] = \left( \frac{1}{1 - \frac{i\nu}{\mu} u} \right)^{\frac{\mu^2}{2\nu t}}$$

Therefore the Log-Characteristic Function could be written as

$$\psi(u, t) = \ln(\Psi(u, t)) = t\psi(u, 1)$$

In other words

$$\mathbb{E}[e^{iuX(t)}] = \Psi(u, t) = \exp(t\psi(u, 1))$$

Using which, the VGSA characteristic function becomes

$$\mathbb{E}[e^{iuX(Y(t))}] = \mathbb{E}[\exp(Y(t)\psi(u, 1))] = \phi(-i\psi(u, 1))$$

with $\phi()$ the CIR characteristic function, namely

$$\phi(u_i) = \mathbb{E}[\exp(iuY_t)] = A(t, u) \exp(B(t, u)y_0)$$

where

$$A(t, u) = \frac{\exp(\kappa^2 \eta t / \lambda^2)}{[\cosh(\gamma t / 2) + \kappa / \gamma \sinh(\gamma t / 2)]^{\frac{\sigma^2}{2\sigma}}}$$

$$B(t, u) = \frac{2\nu u}{\kappa + \gamma \coth(\gamma t / 2)}$$

and

$$\gamma = \sqrt{\kappa^2 - 2\lambda^2}$$

This allows us to determine the VGSA characteristic function, which we can use to calculate options prices via numeric Fourier inversion as described in [48] and [51].

1.11.3 Variance Gamma with Gamma Arrival Rate

For the Variance Gamma with Gamma Arrival Rate (VGG), as before, the Stock Process under the risk-neutral framework is

$$d \ln S_t = (r - q + \omega)dt + X(h(dt); \sigma, \nu, \theta)$$

with $\omega = \frac{1}{\nu} \ln(1 - \theta \nu - \sigma^2 \nu / 2)$ and

$$X(h(dt); \sigma, \nu, \theta) = B(\gamma(h(dt), 1, \nu); \theta, \sigma)$$
and the general Gamma Cumulative Distribution Function for $\gamma(h, \mu, \nu)$ is

$$F(\mu, \nu; h, x) = \frac{1}{\Gamma(\nu/h)} \left( \frac{\mu}{\nu} \right)^{\frac{\nu^2}{h}} \int_0^x e^{-\frac{\mu t}{\nu} t^{\nu-1}} dt$$

and here $h(dt) = dY_t$ with $Y_t$ is also Gamma-distributed

$$dY_t = \gamma(dt, \mu_a, \nu_a)$$

The parameter-set is therefore $\Psi = (\mu_a, \nu_a, \nu, \theta, \sigma)$. 
Chapter 2

The Inference Problem

In applying option pricing models, one always encounters the difficulty that the spot volatility and the structural parameters are unobservable.
- Gurdip Bakshi, Charles Cao & Zhiwu Chen

2.1 Introduction

Regardless of which specific model we are using, it seems that we cannot avoid the issue of calibration. There are two possible sets of data that we can use for estimating the model parameters: options prices and historic stock prices.\(^1\)

Using options prices via a Least Square Estimator (LSE) has the obvious advantage of guaranteeing that we will match the used option market-prices within a certain tolerance. However the availability of option data is typically limited which would force us to use interpolation and extrapolation methods. These data manipulation approaches might deteriorate the quality and the smoothness of our inputs. More importantly, matching a set of plain-vanilla option prices does not necessarily mean that we would obtain the correct price for another more exotic derivative.

Using Stock Prices has the disadvantage of offering no guarantee of matching option prices. However supposing that the model is right, we do have a great quantity of data-input for calibration, which is a powerful argument in favor of this approach.

\(^1\)Recently some researchers have also tried to use Historic Option Prices. See for instance Elliott [93] or Van der Merwe [229].
It is important however to note that in using historic stock prices we are assuming that our time-step $\Delta t$ is small enough so that we are almost in a continuous setting. What is more, we are assuming the validity of the Girsanov theorem which is applicable to a diffusion-based model. This also means we are implicitly assuming that the market price of volatility risk is stable and so are the risk-neutral volatility-drift parameters.

More accurately having for instance a real-world model

$$dv_t = (\omega - \theta v_t)dt + \xi v_t^p dZ_t$$

with $p = 0.5$ corresponding to the Heston model, we know that the risk-neutral volatility-drift parameter is

$$\theta^{(r)} = \theta + \lambda \xi v_t^p$$

As a result, supposing that $\theta^{(r)}$ is a stable (or even constant) parameter, is equivalent to supposing that $\lambda$ the market-price-of-volatility-risk verifies

$$\lambda = \phi v_t^{1-p}$$

with $\phi$ a constant coefficient.

The implication of this assumption for a model with a real-world parameter-set $\Psi = (\omega, \theta, \xi, \rho)$ and a risk-neutral counterpart $\Psi^{(r)} = (\omega^{(r)}, \theta^{(r)}, \xi^{(r)}, \rho^{(r)})$ is

$$\xi = \xi^{(r)}$$
$$\rho = \rho^{(r)}$$
$$\omega = \omega^{(r)}$$
$$\theta = \theta^{(r)} - \phi$$

Let us insist on the fact that the above assumption is valid only for a diffusion-based model. For some non-Gaussian pure-jump models such as VGG, we lose the comparability between the statistical and the risk-neutral parameters. We could instead use the stock-price time-series to determine the statistical density $p(z)$ on the one hand, and use the options-prices to determine the risk-neutral density $q(z)$ on the other; and calculate the ratio $r(z) = p(z)/q(z)$ corresponding to the Radon-Nikodym derivative of the two measures for this model.

$^2$Note that many call the market-price-of-volatility-risk the quantity $\lambda \xi v_t^p$.

$^3$Also as stated by Bakshi, Cao & Chen [20]: When the risk-aversion coefficient of the representative agent is bounded within a reasonable range, the parameters of the true distributions will not differ significantly from their risk-neutral counterparts.

The direct implication of this is

$$\theta \approx \theta^{(r)}$$

More importantly, for daily data we have

$$\Delta t = o(\sqrt{\Delta t})$$

and therefore using either the real-world asset-drift $\mu_S$ or the dividend-adjusted risk-free rate $r - q$ would not make a difference in parameter estimation. Some [10] even ignore the stock drift term altogether.
The central question of this chapter is therefore the *Inference* of the parameters embedded in a stochastic volatility model. The summary of the logical subdivisions of the problem is as follows:

1. **Cross-Sectional vs. Time-Series**: The former uses Options Prices at a given point in time and the latter a series of the underlying prices for a given period. As mentioned earlier, the former provides an estimation of the parameters in the risk-neutral universe and the latter estimation takes place in the statistical universe.

2. **Classical vs. Bayesian**: Using *Time-Series*, one could suppose that there exists an unknown but fixed set of parameters and try to estimate them as closely as possible. This is a Classical (Frequentist) approach. Alternatively one could use a Bayesian approach where the parameters are supposed to be random variables and have their prior distributions that one can update via the observations.

3. **Learning vs. Likelihood-Maximization**: Under the Classical hypothesis, one could either try to estimate the instantaneous variance together with the fixed parameters, which corresponds to a Learning process. A more robust way would be to estimate the Likelihood function and maximize it over all the possible values of the parameters.

4. **Gaussian vs. Non-Gaussian**: In any of the above approaches the stochastic volatility (SV) model could be diffusion-based or not. As we will see further, this will affect the actual estimation methodology. Among the Gaussian SV models we consider, are Heston, GARCH and 3/2. Among the Non-Gaussian ones are Bates, VGS and VGG.

5. **State-Space Representation**: For each of the above approaches and for each SV model, we have a number of ways of choosing a state and represent the instantaneous variance as well as the spot price. Needless to say, a more parsimonious and lower-dimension state is preferable.

6. **Diagnostics & Sampling Distribution**: Once the inference process is finished, one has to verify its accuracy via various Tests. Quantities such as MPE, RMSE, Box-Ljung or $\chi^2$ numbers correspond to some of the possible Diagnostics. Observing the Sampling Distribution over various paths is another way of checking the validity of the inference methodology.

Finally it is worth noting that our entire approach is based upon **Parametric** stochastic volatility models. This model class is more restrictive than the Non or Semi Parametric one, however it has the advantage of offering the possibility of a direct interpretation of the resulting parameters.
2.2 Using Option Prices

Using a set of current vanilla option prices, we can perform an LSE and assess the risk-neutral model parameters. Taking a set of \( J \) strike prices \( K_j \)'s with their corresponding option prices \( C_{mkt}(K_j) \) for a given maturity, we would try to minimize

\[
\sum_{j=1}^{J} (C_{model}(K_j) - C_{mkt}(K_j))^2
\]

The minimization\(^4\) could for example be done via the Direction Set (Powell) method, the Conjugate Gradient (Fletcher-Reeves-Polak-Ribiere) method or the Levenberg-Marquardt (LM) method. We will now briefly describe the Powell optimization algorithm.

2.2.1 Direction Set (Powell) Method

The optimization method we will use later is the Direction Set (Powell) method and does not require any Gradient or Hessian computation.\(^5\) This is a quadratically convergent method producing mutually conjugate directions.

Most multi-dimensional optimization algorithms require a one-dimensional line minimization routine which does the following: Given as input the vectors \( \mathbf{P} \) and \( \mathbf{n} \) and the function \( f \), find the scalar \( \lambda \) that minimizes \( f(\mathbf{P} + \lambda \mathbf{n}) \), then replace \( \mathbf{P} \) with \( \mathbf{P} + \lambda \mathbf{n} \) and \( \mathbf{n} \) with \( \lambda \mathbf{n} \).

The idea would be to take a set of directions that are as non-interfering as possible in order to avoid spoiling one minimization with the subsequent one. This way an interminable cycling through the set of directions will not occur.

This is why we seek conjugate directions. Calling the function to be minimized \( f(\mathbf{x}) \) with \( \mathbf{x} \) a vector of dimension \( N \), we can write the second order Taylor expansion around a particular point \( \mathbf{P} \)

\[
f(\mathbf{x}) \approx f(\mathbf{P}) + \nabla f(\mathbf{P})\mathbf{x} + \frac{1}{2}\mathbf{x}H\mathbf{x}
\]

where \( H_{ij} = \frac{\partial^2 f}{\partial x_i \partial x_j} \) is the Hessian matrix of the function at point \( \mathbf{P} \).

We therefore have for the variation of the gradient \( \delta(\nabla f) = H\delta\mathbf{x} \) and in order

\(^4\)Some consider that this minimization will give more importance to the ATM options and try therefore to correct this by introducing weights in the summation.

\(^5\)There are also Entropy-based techniques as discussed in [16] applied to Local Volatility models, which are different from our parametric models.

This is an important feature when the function to be optimized contains discontinuities.
to have a non-interfering new direction we choose $\mathbf{v}$ such as the motion along $\mathbf{v}$ remains perpendicular to our previous direction $\mathbf{u}$

$$\mathbf{u}\delta(\nabla f) = \mathbf{u}H\mathbf{v} = 0$$

In this case the directions $\mathbf{u}$ and $\mathbf{v}$ are said to be conjugate.

Powell suggests a quadratically convergent method that produces a set of $N$ mutually conjugate directions. The following description is taken from Press [204] where the corresponding source code could be found as well.

1. Initialize the set of directions $\mathbf{u}_i$ to the basis vectors for $i = 1, \ldots, N$
2. Save the starting point as $\mathbf{P}_0$
3. Move $\mathbf{P}_{i-1}$ to the minimum along direction $\mathbf{u}_i$ and call it $\mathbf{P}_i$
4. Set $\mathbf{u}_i$ to $\mathbf{u}_{i+1}$ for $i = 1, \ldots, N - 1$ and set $\mathbf{u}_N$ to $\mathbf{P}_N - \mathbf{P}_0$
5. Move $\mathbf{P}_N$ to the minimum along $\mathbf{u}_N$ and call this point $\mathbf{P}_0$ and go back to step 2

For a quadratic form, $k$ iterations of this algorithm will produce a set of directions whose last $k$ members are mutually conjugate. The idea is to repeat the steps until the function stops decreasing. However, this procedure tends to produce directions that are linearly dependent and therefore provides us with the minimum only over a sub-space. Hence the idea of discarding the direction along which $f$ made the largest decrease. This seems paradoxical, we are dropping our best direction in the new iteration, however this is the best chance of avoiding a build-up of linear dependence.

In what follows we apply the Powell algorithm to SPX options valued via the mixing monte-carlo method.

### 2.2.2 Numeric Tests

We apply the Powell algorithm to SPX Options valued via the Mixing Monte-Carlo method. The optimization is performed across close-to-the-money strike prices as of $t_0 = 2002/05/21$ with the index $S_0 = 1079.88$ and maturities $T = 2002/08/17$, $T = 2002/09/21$, $T = 2002/12/21$, and $T = 2003/03/22$. 
Figure 2.1: S&P500 Volatility Surface as of 20020521 with Index=1079.88 USD. The surface will be used for fitting via the Direction Set (Powell) Optimization algorithm applied to a Square-Root model implemented with a one-factor Monte-Carlo Mixing method.
Figure 2.2: Mixing Monte-Carlo Simulation with the Square-Root Model for SPX on 20020521 with Index=1079.88 USD, Maturity 20020817

Powell (Direction Set) Optimization method was used for Least-Square Calibration

Optimal Parameters $\hat{\omega} = 0.081575$, $\hat{\theta} = 3.308023$, $\hat{\xi} = 0.268151$, $\hat{\rho} = -0.999999$
Figure 2.3: Mixing Monte-Carlo Simulation with the Square-Root Model for SPX on 20020521 with Index=1079.88 USD, Maturity 20020921
Powell (Direction Set) Optimization method was used for Least-Square Calibration
Optimal Parameters $\hat{\omega} = 0.108359, \hat{\theta} = 3.798900, \hat{\xi} = 0.242820, \hat{\rho} = -0.999830$
Figure 2.4: Mixing Monte-Carlo Simulation with the Square-Root Model for SPX on 20020521 with Index=1079.88 USD, Maturity 20021221
Powell (Direction Set) Optimization method was used for Least-Square Calibration
Optimal Parameters $\hat{\omega} = 0.126519, \hat{\theta} = 3.473910, \hat{\xi} = 0.222532, \hat{\rho} = -0.999991$
Figure 2.5: Mixing Monte-Carlo Simulation with the Square-Root Model for SPX on 20020521 with Index=1079.88 USD, Maturity 20030322
Powell (Direction Set) Optimization method was used for Least-Square Calibration
Optimal Parameters $\hat{\omega} = 0.138687, \hat{\theta} = 3.497779, \hat{\xi} = 0.180010, \hat{\rho} = -1.000000$
As we see in table 2.1, the estimated parameters are fairly stable for different maturities and therefore the stochastic volatility model seems to be fairly time-homogeneous.

<table>
<thead>
<tr>
<th>T</th>
<th>ω</th>
<th>θ</th>
<th>ξ</th>
<th>ρ</th>
</tr>
</thead>
<tbody>
<tr>
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<td>0.081575</td>
<td>3.308023</td>
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<tr>
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<td>0.138687</td>
<td>3.497779</td>
<td>0.180010</td>
<td>-1.000000</td>
</tr>
</tbody>
</table>

Table 2.1: The estimation is performed for SPX on t=20020521 with Index=1079.88 USD for different maturities T.

2.2.3 The Distribution of the Errors

Since the parameter-set Ψ contains only a few elements and we can have many options prices, it is clear that the matching of the model and market prices is not perfect. Hence the idea of observing the distribution of the errors

$$C_{mkt}(K_j) = C_{model}(K_j, \hat{\Psi}) \exp\left\{ -\frac{1}{2} \Upsilon^2 + \Upsilon N^{(j)}(0,1) \right\}$$

with $1 \leq j \leq J$ and $\Upsilon$ the error standard-deviation, and $\hat{\Psi}$ the optimal parameter-set. As usual $N(0,1)$ is the standard Normal distribution.

Note that our previously discussed LSE approach is not exactly equivalent to the maximization of a Likelihood function based upon the above distribution, since the latter would correspond to the minimization of the sum of the squared Log returns.

A good Bias test would be to check for the predictability of the errors. For this, one could run a regression of the error

$$e_j = C_{mkt}(K_j) - C_{model}(K_j, \hat{\Psi})$$

upon a few factors corresponding for instance to moneyness or maturity. A low $R^2$ for the regression would prove that the model errors are not predictable and there is no major bias. For a detailed study, see [182] for instance.
2.3 Using Stock Prices

2.3.1 The Likelihood Function
If as in the previous section, we use European Options with a given maturity \( T \) and with different strike prices, then we will be estimating
\[
q(S_T|S_0; \Psi)
\]
which corresponds to the risk-neutral density, given a known current stock price \( S_0 \) and given a constant parameter-set \( \Psi \). As discussed, the Least-Squares Estimation (LSE) is used to find the best guess for the unknown ideal parameter-set.

On the other hand if we use a time-series of stock-prices \( (S_t)_{0 \leq t \leq T} \) we would be dealing with the \textit{joint} probability
\[
p(S_1, \ldots, S_T|S_0; \Psi)
\]
which we can rewrite as
\[
p(S_1, \ldots, S_T|S_0; \Psi) = \prod_{t=1}^{T} p(S_t|S_{t-1}, \ldots, S_0; \Psi)
\]
It is the above joint probability that is commonly referred to as the Likelihood Function \( L_{0:T}(\Psi) \). Maximizing the Likelihood over the parameter-set \( \Psi \) would provide us with the best parameter-set for the statistical density \( p(S_T|S_0; \Psi) \).

Note that we are using a Classical (Frequentist) approach where we assume that the parameters are unknown but fixed over \([0, T]\). In other words we would be dealing with the \textit{same} parameter-set for any of the \( p(S_t|S_{t-1}, \ldots, S_0; \Psi) \) with \( 1 \leq t \leq T \).

It is often convenient to work with the Log of the Likelihood function since this will produce a sum
\[
\ln L_{0:T}(\Psi) = \sum_{t=1}^{T} \ln p(S_t|S_{t-1}, \ldots, S_0; \Psi)
\]

The Justification for the MLE
As explained for instance in [100], one justification of the maximization of the (Log) Likelihood function, comes from the \textit{Kullback-Leibler} (KL) distance. The KL distance is defined as\(^6\)
\[
d(p^*, p) = \int p^*(x) (\ln p^*(x) - \ln p(x)) \, dx
\]
\(^6\)Hereafter when the bounds are not specified, the integral is taken on the entire space of the integrand argument.
where $p^*(x)$ is the ideal density, while $p(x)$ the density under estimation.

We can write

$$d(p^*, p) = \mathbf{E}^* \ln (p^*(x)/p(x))$$

Note that using the Jensen (Log convexity) inequality

$$d(p^*, p) = -\mathbf{E}^* \ln (p(x)/p^*(x)) \geq -\ln (\mathbf{E}^*(p(x)/p^*(x)))$$

so

$$d(p^*, p) \geq -\ln \int p^*(x)p(x)/p^*(x)dx = 0$$

and $d(p, p^*) = 0$ if and only if $p = p^*$, which confirms that $d(.,.)$ is a distance.

Now minimizing $d(p, p^*)$ over $p()$ would be equivalent to minimizing the term

$$-\int p^*(x) \ln p(x) dx$$

since the rest of the expression depends on $p^*(.)$ only.

This latter expression could be written in the discrete framework, having $T$ observations $S_1, ..., S_T$ as

$$-\sum_{t=1}^{T} \ln p(S_t)$$

since the observations are by assumption distributed according to the ideal $p^*(.)$.

This justifies our maximizing

$$\prod_{t=1}^{T} p(S_t)$$

Note that in a pure parameter estimation, this would be the MLE approach. However the minimization of the KL distance is more general and can allow for Model Identification.

MLE has many desirable asymptotic attributes as explained for example in [127]. Indeed ML estimators are consistent and converge to the right parameter-set as the number of observations increases. They actually reach the lower bound for the error, referred to as the Cramer-Rao bound which corresponds to the inverse of the Fisher Information matrix.

Calling the first derivative of the Log-likelihood the score function

$$h(\Psi) = \frac{\partial \ln L_{0:T}(\Psi)}{\partial \Psi}$$

it is known that MLE could be interpreted as a special case of the General Method of Moments (GMM) where the moment $g(\Psi)$ such that

$$\mathbf{E}[g(\Psi)] = 0$$
is simply taken to be the above score function. In need we would then have
\[ E[h(\Psi)] = \int \frac{\partial \ln L_{0:T}(\Psi)}{\partial \Psi} L_{0:T}(\Psi) d_{0:T} = 0 \]
which means that
\[ \int \frac{\partial L_{0:T}(\Psi)}{\partial \Psi} d_{0:T} = 0 \]
as previously discussed in the MLE.

Note that taking the derivative of the above with respect to the parameter-set (using one-dimensional notations for simplicity)
\[ \int \frac{\partial}{\partial \Psi} (h(\Psi)L_{0:T}(\Psi)) d_{0:T} = 0 \]
which will give us
\[ \int \frac{\partial^2 \ln L_{0:T}(\Psi)}{\partial \Psi^2} L_{0:T}(\Psi) d_{0:T} = -\int \frac{\partial L_{0:T}(\Psi)}{L_{0:T}(\Psi)} \frac{\partial L_{0:T}(\Psi)}{\partial \Psi} d_{0:T} = -\int \left( \frac{\partial \ln L_{0:T}(\Psi)}{\partial \Psi} \right)^2 L_{0:T}(\Psi) d_{0:T} \]
meaning that
\[ J = -E \left[ \frac{\partial^2 \ln L_{0:T}(\Psi)}{\partial \Psi^2} \right] = E \left[ \left( \frac{\partial \ln L_{0:T}(\Psi)}{\partial \Psi} \right)^2 \right] \]
which is referred to as the Information Matrix Identity.

As previously stated, asymptotically we have for the optimal parameter-set \( \hat{\Psi} \) and the ideal \( \Psi^\ast \)
\[ \hat{\Psi} - \Psi^\ast \sim \mathcal{N} (0, J^{-1}) \]

**Likelihood Evaluation and Filtering**

For GARCH models the Likelihood is known under an integrated form. Indeed calling \( u_t \) the mean-adjusted stock return, \( v_t \) the variance and \( (B_t) \) a Gaussian sequence, we have for any GARCH model
\[ u_t = h(v_t, B_t) \]
and
\[ v_t = f(v_{t-1}, u_{t-1}; \Psi) \]
where $f()$ and $h()$ are two deterministic functions. This will allow us to directly determine and optimize\(^7\)

$$L_{1:T}(\Psi) \propto - \sum_{t=1}^{T} \ln(v_t) + \frac{u_t^2}{v_t}$$

This is possible because GARCH models have one source of randomness and there is a time-shift between the variance and the spot equations.

Unlike GARCH, most Stochastic Volatility models have two (non-perfectly correlated) sources of randomness ($B_t$) and ($Z_t$) and have equations of the form

$$u_t = h(v_t, B_t)$$
$$v_t = f(v_{t-1}, Z_t; \Psi)$$

which means that the Likelihood function is not directly known under an integrated form, and we need Filtering techniques for its estimation and optimization.

Another justification for Filtering is its application to Parameter Learning. As we shall see, in this approach we use the joint distribution of the hidden state and the parameters. In order to obtain the optimal value of the hidden state $v_t$ given all the observations $z_{1:t}$ we need to use a Filter.

### 2.3.2 Filtering

The idea here is to use the Filtering Theory for the estimation of Stochastic Volatility model parameters. What we are trying to do is to find the probability density function (pdf) corresponding to a state $x_k$ at time step $k$ given all the observations $z_{1:k}$ up to that time.

Looking for the pdf $p(x_k|z_{1:k})$ we can proceed in two stages:

1) First we can write the Time Update iteration by applying the Chapman-Kolmogorov equation\(^8\)

$$p(x_k|z_{1:k-1}) = \int p(x_k|x_{k-1}, z_{1:k-1})p(x_{k-1}|z_{1:k-1})dx_{k-1} = \int p(x_k|x_{k-1})p(x_{k-1}|z_{1:k-1})dx_{k-1}$$

by using the Markov property.

---

\(^7\)We generally drop constant terms in the likelihood function since they do not affect the optimal arguments. Hence the notation $L_{1:T}(\Psi) \propto ...$

\(^8\)See Shreve [218] for instance.
2) Following this, for the Measurement Update we use the Bayes rule
\[ p(x_k|z_{1:k}) = \frac{p(z_k|x_k)p(x_k|z_{1:k-1})}{p(z_k|z_{1:k-1})} \]
where the denominator \( p(z_k|z_{1:k-1}) \) could be written as
\[ p(z_k|z_{1:k-1}) = \int p(z_k|x_k)p(x_k|z_{1:k-1})dx_k \]
and corresponds to the Likelihood Function for the time-step \( k \).

**Proof:**
Indeed we have
\[
\begin{align*}
p(x_k|z_{1:k}) &= \frac{p(z_{1:k}|x_k)p(x_k)}{p(z_{1:k})} \\
&= \frac{p(z_k,z_{1:k-1}|x_k)p(x_k)}{p(z_k,z_{1:k-1})} \\
&= \frac{p(z_k|z_{1:k-1},x_k)p(z_{1:k-1}|x_k)p(x_k)}{p(z_k|z_{1:k-1})p(z_{1:k-1})} \\
&= \frac{p(z_k|z_{1:k-1},x_k)p(x_k|z_{1:k-1})p(z_{1:k-1})p(x_k)}{p(z_k|z_{1:k-1})p(z_{1:k-1})p(x_k)} \\
&= \frac{p(z_k|x_k)p(x_k|z_{1:k-1})}{p(z_k|z_{1:k-1})}
\end{align*}
\]
which proves the claimed result. \((QED)\)

Note that we use the fact that at time-step \( k \) the value of \( z_{1:k} \) is perfectly known.

The Kalman Filter (detailed below) is a special case where the distributions are Normal and could be written as
\[
p(x_k|z_{k-1}) = \mathcal{N}(\hat{x}_k^- , P^-) \\
p(x_k|z_k) = \mathcal{N}(\hat{x}_k , P_k)
\]
Indeed in the special Gaussian case, each distribution could be entirely characterized via its first two moments. However, it is important to remember that the Kalman Filter (KF) is optimal in the Gaussian linear case. In the Non-Linear case, it will indeed always be sub-optimal.
**Interpretation of the Kalman Gain**

The basic idea behind the KF is the following observation. Having \( x \) a Normally distributed random-variable with a mean \( m_x \) and variance \( S_{xx} \), and \( z \) a Normally distributed random-variable with a mean \( m_z \) and variance \( S_{zz} \), and having \( S_{xz} = S_{zx} \) the covariance between \( x \) and \( z \), the conditional distribution of \( x|z \) is also normal with

\[
m_{x|z} = m_x + K(z - m_z)
\]

with

\[
K = S_{xz}S_{zz}^{-1}
\]

Interpreting \( x \) as the hidden-state and \( z \) as the observation, the above matrix \( K \) would correspond to the Kalman filter in the linear case.

We also have

\[
S_{x|z} = S_{xx} - KS_{xz}
\]

An alternative interpretation of the Kalman Filter could be based upon Linear Regression. Indeed, if we knew the time-series of \((z_k)\) and \((x_k)\), then the regression could be written as

\[
x_k = \beta z_k + \alpha + \epsilon_k
\]

with \( \beta \) the slope, \( \alpha \) the intercept and \((\epsilon_k)\) the residuals. It is known that under a Least Squares regression, we have

\[
\beta = S_{xz}S_{zz}^{-1}
\]

which again is the expression for the Kalman Gain.

We now will describe various non-linear extensions of the Kalman Filter.
2.3.3 The Simple and Extended Kalman Filters

The first algorithms we choose here are the Simple and Extended Kalman Filters\(^9\) due to their well-known flexibility and ease of implementation. The simple or traditional Kalman Filter (KF) applies to linear Gaussian cases, while the Extended KF (EKF) could be used for nonlinear Gaussian cases via a first-order linearization. We shall therefore describe EKF and consider the simple KF as a special case.

In order to clarify the notations, let us briefly rewrite the EKF equations. Given a dynamic process \(x_k\) following a possibly non-linear transition equation

\[
x_k = f(x_{k-1}, w_k)
\]  
(2.1)

we suppose we have a measurement \(z_k\) via a possibly non-linear observation equation

\[
z_k = h(x_k, u_k)
\]  
(2.2)

where \(w_k\) and \(u_k\) are two mutually-uncorrelated sequences of temporally-uncorrelated Normal random-variables with zero means and covariance matrices \(Q_k, R_k\) respectively.\(^{10}\) Moreover, \(w_k\) is uncorrelated with \(x_{k-1}\) and \(u_k\) uncorrelated with \(x_k\).

We define the \textit{linear a priori} process estimate as

\[
\hat{x}_k^- = E[x_k]
\]  
(2.3)

which is the estimation at time step \(k - 1\) prior to measurement. Similarly, we define the \textit{linear a posteriori} estimate

\[
\hat{x}_k = E[x_k|z_k]
\]  
(2.4)

which is the estimation at time step \(k\) after the measurement.

We also have the corresponding estimation errors \(e_k^- = x_k - \hat{x}_k^-\) and \(e_k = x_k - \hat{x}_k\) and the estimate error covariances

\[
P_k^- = E[e_k^- e_k^-']
\]  
(2.5)

\[
P_k = E[e_k e_k']
\]  
(2.6)

where the superscript \(t\) corresponds to the transpose operator.

We now define the Jacobian matrices of \(f\) with respect to the system process and the system noise as \(A_k\) and \(W_k\) respectively. Similarly, we define the gradient matrices of \(h\) with respect to the system process and the measurement noise as \(H_k\) and \(U_k\) respectively.

\(^9\)For a description see for instance Welch [233] or Harvey [129].

\(^{10}\)Some prefer to write \(x_k = f(x_{k-1}, w_{k-1})\). Needless to say, the two notations are equivalent.
More accurately, for every row $i$ and column $j$ we have
\[
A_{ij} = \partial f_i / \partial x_j(\hat{x}_{k-1}, 0) \quad W_{ij} = \partial f_i / \partial w_j(\hat{x}_{k-1}, 0)
\]
\[
H_{ij} = \partial h_i / \partial x_j(\hat{x}_k, 0) \quad U_{ij} = \partial h_i / \partial u_j(\hat{x}_k, 0)
\]

We therefore have the following Time Update equations
\[
\hat{x}_k = f(\hat{x}_{k-1}, 0) \tag{2.7}
\]
and
\[
P_k^- = A_k P_{k-1} A_k^t + W_k Q_{k-1} W_k^t \tag{2.8}
\]

We define the Kalman gain as the matrix $K_k$ used in the Measurement Update equations
\[
\hat{x}_k = \hat{x}_k^- + K_k (z_k - h(\hat{x}_k^-, 0)) \tag{2.9}
\]
and
\[
P_k^- = (I - K_k H_k) P_k^- \tag{2.10}
\]
where $I$ represents the Identity matrix.

The optimal Kalman gain corresponds to the mean of the conditional distribution of $x_k$ upon the observation $z_k$ or equivalently, the matrix that would minimize the mean square error $P_k$ within the class of linear estimators. This optimal gain is
\[
K_k = P_k^- H_k^t (H_k P_k^- H_k^t + U_k R_k U_k^t)^{-1} \tag{2.11}
\]

The above equations complete the Kalman Filter algorithm.

Another Interpretation of the Kalman Gain

Note that an easy way to observe that $K_k$ minimizes the a posteriori error covariance $P_k$ is to consider the one-dimensional linear case
\[
\hat{x}_k = \hat{x}_k^- + K_k (z_k - H_k \hat{x}_k^-) = \hat{x}_k^- + K_k (z_k - H_k x_k + H_k e_k^-)
\]
so
\[
e_k = x_k - \hat{x}_k = e_k^- - K_k (u_k + H_k e_k^-)
\]
Therefore
\[
P_k = E(e_k^2) = P_k^- + K_k^2 (R_k + H_k^2 P_k^- + 0) - 2K_k H_k P_k^-
\]
and taking the derivative with respect to $K_k$ and setting it to zero, we get

$$K_k = \frac{P_k^- H_k}{H_k^2 P_k^- + R_k}$$

which is the one-dimensional expression for the linear Kalman Gain.

**Residuals, MPE and RMSE**

In what follows we shall call the estimated observations $\hat{z}_k^-$. For the Simple and Extended Kalman Filters we have

$$\hat{z}_k^- = h(\hat{x}_k^-, 0)$$

The *residuals* are the observation errors, defined as

$$\tilde{z}_k = z_k - \hat{z}_k^-$$

Needless to say the smaller these residuals, the higher the quality of the Filter. Therefore to measure the performance, we define the Mean Price Error (MPE) and Root Mean Square Error (RMSE) as the mean and standard-deviation of the residuals

$$MPE = \frac{1}{N} \sum_{k=1}^{N} \tilde{z}_k$$

$$RMSE = \sqrt{\frac{1}{N} \sum_{k=1}^{N} (\tilde{z}_k - MPE)^2}$$
2.3.4 The Unscented Kalman Filter

Recently Julier and Uhlmann [166] proposed a new extension of the Kalman Filter to Nonlinear systems, completely different from the EKF. They argue that EKF could be difficult to implement and more importantly difficult to tune and only reliable for systems which are almost linear within the update intervals. The new method called the Unscented Kalman Filter (UKF) will calculate the mean to a higher order of accuracy than the EKF and the covariance to the same order of accuracy.

Unlike the EKF, this method does not require any Jacobian calculation since it does not approximate the nonlinear functions of the process and the observation. Therefore it uses the true nonlinear models but approximates the distribution of the state random variable $x_k$ by applying an Unscented Transformation to it. As we will see below, we construct a set of Sigma Points which capture the mean and covariance of the original distribution and when propagated through the true non-linear system, capture the posterior mean and covariance accurately to the third order.

Similarly to the EKF, we start with an initial choice for the state vector $\hat{x}_0 = \mathbf{E}[x_0]$ and its Covariance Matrix $P_0 = \mathbf{E}[(x_0 - \hat{x}_0)(x_0 - \hat{x}_0)^T]$.

We then concatenate the space vector with the system noise and the observation noise\footnote{This space augmentation will not be necessary if we have additive noises as in $x_k = f(x_{k-1}) + w_{k-1}$ and $z_k = h(x_k) + u_k$} and create an augmented state vector for each step $k$ greater than one

$$x_{a,k-1}^a = \begin{pmatrix} x_{k-1} \\ w_{k-1} \\ u_{k-1} \end{pmatrix}$$

and therefore

$$\hat{x}_{a,k-1}^a = \begin{pmatrix} \hat{x}_{k-1} \\ 0 \\ 0 \end{pmatrix}$$

and

$$P_{a,k-1}^a = \begin{pmatrix} P_{k-1} & P_{xw} (k-1|k-1) & 0 \\ P_{xw} (k-1|k-1) & P_{ww} (k-1|k-1) & 0 \\ 0 & 0 & P_{uu} (k-1|k-1) \end{pmatrix}$$

for each iteration $k$. The augmented state will therefore have a dimension $n_a = n_x + n_w + n_u$.

We then need to calculate the corresponding Sigma Points through the Unscented Transformation:

$$\chi_{k-1}^a(0) = \hat{x}_{k-1}^a$$

For $i = 1, \ldots, n_a$

$$\chi_{k-1}^a(i) = \hat{x}_{k-1}^a + (\sqrt{(n_a + \lambda)P_{a,k-1}^a})_i$$
and for $i = n_a + 1, \ldots, 2n_a$

$$
\chi_{k-1}^a(i) = \tilde{x}_{k-1}^a - (\sqrt{(n_a + \lambda)P_{k-1}^{a}})_{i-n_a}
$$

where the above subscripts $i$ and $i-n_a$ correspond to the $i^{th}$ and $i-n_a^{th}$ columns of the square-root matrix.\footnote{The square-root matrix is calculated via Singular Value Decomposition (SVD) and Cholesky factorization \cite{204}. In case $P_{k-1}^a$ is not Positive-Definite, one could for example use a truncation procedure.} This prepares us for the Time Update and the Measurement Update equations, similarly to the EKF.

The \textit{Time Update} equations are

$$
\chi_{k|k-1}(i) = f(\chi_{k-1}^x(i), \chi_{k-1}^w(i))
$$

for $i = 0, \ldots, 2n_a + 1$ and

$$
\tilde{x}_k^- = \sum_{i=0}^{2n_a} W_i^{(m)} \chi_{k|k-1}(i)
$$

and

$$
P_k^- = \sum_{i=0}^{2n_a} W_i^{(c)} (\chi_{k|k-1}(i) - \tilde{x}_k^-)(\chi_{k|k-1}(i) - \tilde{x}_k^-)^t
$$

where the superscripts $x$ and $w$ respectively correspond to the state and system-noise portions of the augmented state.

The $W_i^{(m)}$ and $W_i^{(c)}$ weights are defined as

$$
W_0^{(m)} = \frac{\lambda}{n_a + \lambda}
$$

and

$$
W_0^{(c)} = \frac{\lambda}{n_a + \lambda} + (1 - \alpha^2 + \beta)
$$

and for $i = 1, \ldots, 2n_a$

$$
W_i^{(m)} = W_i^{(c)} = \frac{1}{2(n_a + \lambda)}
$$

The scaling parameters $\alpha$, $\beta$, $\kappa$ and $\lambda = \alpha^2(n_a + \kappa) - n_a$ will be chosen for tuning.

We also define within the Time Update equations

$$
Z_{k|k-1}(i) = h(\chi_{k|k-1}(i), \chi_{k-1}^u(i))
$$

and

$$
\tilde{z}_k^- = \sum_{i=0}^{2n_a} W_i^{(m)} Z_{k|k-1}(i)
$$
where the superscript $u$ corresponds to the observation-noise portion of the augmented state.

As for the *Measurement Update* equations, we have

$$
P_{z_k z_k} = \sum_{i=0}^{2n_c} W_i^{(c)} (Z_{k|k-1}(i) - \hat{z}_k^-)(Z_{k|k-1}(i) - \hat{z}_k^-)^t
$$

and

$$
P_{x_k z_k} = \sum_{i=0}^{2n_c} W_i^{(c)} (\chi_{k|k-1}(i) - \hat{x}_k^-)(Z_{k|k-1}(i) - \hat{z}_k^-)^t
$$

which gives us the Kalman Gain

$$
K_k = P_{x_k z_k} P_{z_k z_k}^{-1}
$$

and we get as before

$$
\hat{x}_k = \hat{x}_k^- + K_k (z_k - \hat{z}_k^-)
$$

where again $z_k$ is the observation at time (iteration) $k$.

Also, we have

$$
P_k = P_k^- - K_k P_{z_k z_k} K_k^t
$$

which completes the Measurement Update Equations.
2.3.5 Kushner’s Non-Linear Filter

It would be instructive to compare this algorithm to the nonlinear filtering algorithm based on an approximation of the conditional distribution by Kushner et al. [174]. In this approach, the authors suggest using a Gaussian Quadrature in order to calculate the integral at the measurement update (or the time update) step.\(^\text{13}\)

As the Kushner paper indicates, having an \(N\)-dimensional normal random-variable \(X = \mathcal{N}(m, P)\) with \(m\) and \(P\) the corresponding mean and covariance, for a polynomial \(G\) of degree \(2M - 1\) we can write\(^\text{14}\)

\[
E[G(X)] = \frac{1}{(2\pi)^{\frac{N}{2}}|P|^\frac{1}{2}} \int G(y) \exp\left[-\frac{(y - m)^tP^{-1}(y - m)}{2}\right] dy
\]

which is equal to

\[
E[G(X)] = \sum_{i_1=1}^{M} \ldots \sum_{i_N=1}^{M} w_{i_1} \ldots w_{i_N} G(m + \sqrt{P}\zeta)
\]

where \(\zeta^t = (\zeta_{i_1}, \ldots, \zeta_{i_N})\) is the vector of the Gauss-Hermite roots of order \(M\) and \(w_{i_1}, \ldots, w_{i_N}\) are the corresponding weights.

Note that even if both Kushner’s NLF and UKF use Gaussian Quadratures, UKF only uses \(2N + 1\) sigma points, while NLF needs \(MN\) points for the integral computation.

Kushner and Budhiraja suggest using this technique primarily for the measurement update (filtering) step. They claim that provided this step is properly implemented, the time update (prediction) step can be carried out via a linearization similarly to the EKF.

Details of the Kushner algorithm

Let us use the same notations as for the UKF algorithm. We therefore have the augmented state \(x_{k-1}^a\) and its covariance \(P_{k-1}^a\) as before. Here, for a Quadrature order of \(M\) on an \(N\)-dimensional variable, the sigma-points are defined for \(j = 1, ..., N\) and \(i_j = 1, ..., M\) as

\[
\chi_{k-1}^a(i_1, ..., i_N) = x_{k-1}^a + \sqrt{P_{k-1}^a}\zeta(i_1, ..., i_N)
\]

where the square-root here corresponds to the Cholesky factorization, and again \(\zeta(i_1, ..., i_N)[j] = \zeta_{i_j}\) for each \(j\) between 1 and the dimension \(N\) and each \(i_j\)

\(^{13}\)The analogy between Kushner’s Nonlinear Filter and the Unscented Kalman Filter, has already been studied in Ito & Xiong [151].

\(^{14}\)A description of the Gaussian Quadrature could be found in Press et al. [204].
between 1 and the Quadrature order $M$.
Similarly to the UKF, we have the Time Update equations

$$
\chi_{k|k-1}(i_1, \ldots, i_N) = f(\chi_{k-1}^x(i_1, \ldots, i_N), \chi_{k-1}^w(i_1, \ldots, i_N))
$$

but now

$$
\hat{x}_k^- = \sum_{i_1=1}^{M} \cdots \sum_{i_N=1}^{M} w_{i_1} \cdots w_{i_N} \chi_{k|k-1}(i_1, \ldots, i_N)
$$

and

$$
P_k^- = \sum_{i_1=1}^{M} \cdots \sum_{i_N=1}^{M} w_{i_1} \cdots w_{i_N} (\chi_{k|k-1}(i_1, \ldots, i_N) - \hat{x}_k^-)(\chi_{k|k-1}(i_1, \ldots, i_N) - \hat{x}_k^-)^t
$$

Again, we have

$$
Z_{k|k-1}(i_1, \ldots, i_N) = h(\chi_{k|k-1}(i_1, \ldots, i_N), \chi_{k-1}^u(i_1, \ldots, i_N))
$$

and

$$
\hat{z}_k^- = \sum_{i_1=1}^{M} \cdots \sum_{i_N=1}^{M} w_{i_1} \cdots w_{i_N} Z_{k|k-1}(i_1, \ldots, i_N)
$$

Therefore the Measurement Update Equations will be

$$
P_{z_k z_k} = \sum_{i_1=1}^{M} \cdots \sum_{i_N=1}^{M} w_{i_1} \cdots w_{i_N} (Z_{k|k-1}(i_1, \ldots, i_N) - \hat{z}_k^-)(Z_{k|k-1}(i_1, \ldots, i_N) - \hat{z}_k^-)^t
$$

and

$$
P_{x_k z_k} = \sum_{i_1=1}^{M} \cdots \sum_{i_N=1}^{M} w_{i_1} \cdots w_{i_N} (\chi_{k|k-1}(i_1, \ldots, i_N) - \hat{x}_k^-)(Z_{k|k-1}(i_1, \ldots, i_N) - \hat{z}_k^-)^t
$$

which gives us the Kalman Gain

$$
K_k = P_{x_k z_k} P_{z_k z_k}^{-1}
$$

and we get as before

$$
\hat{x}_k = \hat{x}_k^- + K_k(z_k - \hat{z}_k^-)
$$

where again $z_k$ is the observation at time (iteration) $k$.
Also, we have

$$
P_k = P_k^- - K_k P_{z_k z_k} K_k^t
$$

which completes the Measurement Update Equations.

When $N = 1$ and $\lambda = 2$, the numeric integration in the UKF will correspond to a Gauss-Hermite Quadrature of order $M = 3$. However in the UKF we can tune the filter and reduce the higher term errors via the previously mentioned
parameters $\alpha$ and $\beta$.

Note that when $h(x, u)$ is strongly nonlinear, the Gauss Hermite integration is not efficient for evaluating the moments of the measurement update equation, since the term $p(z_k|x_k)$ contains the exponent $z_k - h(x_k, u_k)$. The iterative methods based on the idea of importance sampling proposed in [174] correct this problem at the price of a strong increase in computation time. As suggested in [151], one way to avoid this integration would be to make the additional hypothesis that $x_k, h(x_k, u_k)|z_{1:k-1}$ is Gaussian.
2.3.6 Parameter Learning

One important issue to realize is that the Kalman Filter can be used either for State Estimation (Filtering) or for Parameter Estimation (Machine Learning). When we have both State Estimation and Parameter Estimation we are dealing with a Dual or Joint Estimation. This latter case is the one concerning us since we are estimating the state volatility as well as the model parameters.

As explained in Haykin’s book [133], in a Dual Filter we separate the state vector from the parameters and we apply two intertwined filters to them. On the other hand in a Joint Filter, we concatenate the state vector and the parameters and apply one filter to this augmented state. Note that in the Dual Filter we need to compute recurrent derivatives with respect to parameters, while in a Joint Filter no such step is needed.

It is possible to interpret the Joint Filter in the following way. In a regular filter, i.e. filtering of the state $x_k$ for a fixed set of parameters $\Psi_0$, we are maximizing the conditional density

$$p(x_{1:k}|z_{1:k}, \Psi_0)$$

and as we said, to do that we write

$$p(x_{1:k}|z_{1:k}, \Psi_0) = \frac{p(z_{1:k}|x_{1:k}, \Psi_0)p(x_{1:k}|\Psi_0)}{p(z_{1:k}|\Psi_0)}$$

so we maximize the above with respect to the state $x_k$ for a given set of parameters.

This means that the optimal state $\hat{x}_{1:k}$ for a given parameter-set is given by

$$\hat{x}_{1:k} = \arg\max[p(z_{1:k}, x_{1:k}|\Psi_0)]$$

As we will see, in an MLE approach we use this optimal state filtering for each iteration of the likelihood-maximization over the parameter-set $\Psi$.

In a Joint Filter on the other hand, we are directly optimizing the joint conditional density

$$p(x_{1:k}, \Psi|z_{1:k})$$

which we can also write as

$$p(x_{1:k}, \Psi|z_{1:k}) = \frac{p(z_{1:k}|x_{1:k}, \Psi)p(x_{1:k}|\Psi)p(\Psi)}{p(z_{1:k})}$$

given that the denominator is functionally independent of $x_{1:k}$ and $\Psi$, and given that $p(\Psi)$ contains no prior information, the maximization will be upon

$$p(z_{1:k}|x_{1:k}, \Psi)p(x_{1:k}|\Psi) = p(z_{1:k}, x_{1:k}|\Psi)$$

Again we are in a Frequentist framework, not Bayesian.
In other words in a Joint Filter, the optimal state $\hat{x}_{1:k}$ and parameter-set $\hat{\Psi}$ are found by writing

$$(\hat{x}_{1:k}, \hat{\Psi}) = \arg\max [p(z_{1:k}, x_{1:k} | \Psi)]$$

In what follows we apply the Joint EKF methodology to a few examples.

**An Illustration**

Before using this technique for the Stochastic Volatility model, let us take a simple example where

$$\xi_k = \xi_{k-1} + \pi + 0.10 w_k$$

and

$$z_k = \xi_k + 0.10 u_k$$

where $\pi \approx 3.14159$ and $w_k, u_k$ are independent Gaussian random variables.

The linear state-space system could be written as

$$x_k = \begin{pmatrix} \xi_k \\ \pi_k \end{pmatrix} = \begin{pmatrix} 1 & 1 \\ 0 & 1 \end{pmatrix} x_{k-1} + 0.10 \begin{pmatrix} w_k \\ 0 \end{pmatrix}$$

and

$$z_k = \begin{pmatrix} 1 & 0 \end{pmatrix} x_k + 0.10 u_k$$

We choose the initial values $\xi_0 = z_0 = 0$ and $\pi_0 = 1.0$. We also take $Q = 0.1 I_2$ and $R = 0.10$.

Applying the Kalman Filter to an artificially generated data-set, we plot the resulting $\pi_k$ in figure 2.6. As we can see, the parameter converges very quickly to its true value.

Even if we associated a noise of 0.10 to the constant parameter $\pi$ we can see that for 5000 observations, taking the mean of the filtered state between observations 20 and 5000 we get

$$\hat{\pi} = 3.141390488$$

which is very close to the value 3.14159 used in data generation process.
Figure 2.6: A simple example for the Joint Filter. The convergence towards the constant parameter $\pi$ happens after a few iterations.
Joint Filtering Examples

After going through this simple example, we now apply the JF technique to our Stochastic Volatility problem. We shall study a few examples in order to find the best state space representation.

**Example 1** Our first example would be the Square-Root Stochastic Volatility model

\[
\ln S_k = \ln S_{k-1} + (\mu_S - \frac{1}{2} v_{k-1}) \Delta t + \sqrt{v_{k-1}} \sqrt{\Delta t} B_{k-1}
\]

\[
v_k = v_{k-1} + (\omega - \theta v_{k-1}) \Delta t + \xi \sqrt{v_{k-1} \Delta t} Z_{k-1}
\]

To simplify we suppose that the value of \(\mu_S\) is known.

We can now define the state variable\(^{16}\)

\[
x_k = \begin{pmatrix}
\ln S_k \\
v_k \\
\omega \\
\theta \\
\xi \\
\rho
\end{pmatrix}
\]

and the system noise

\[
w_k = \begin{pmatrix}
B_k \\
Z_k
\end{pmatrix}
\]

with its covariance matrix

\[
Q_k = \begin{pmatrix}
1 & \rho \\
\rho & 1
\end{pmatrix}
\]

and therefore

\[
x_k = f(x_{k-1}, w_{k-1}) = \begin{pmatrix}
\ln S_{k-1} + (\mu_S - \frac{1}{2} v_{k-1}) \Delta t + \sqrt{v_{k-1}} \sqrt{\Delta t} B_{k-1} \\
v_{k-1} + (\omega - \theta v_{k-1}) \Delta t + \xi \sqrt{v_{k-1} \Delta t} Z_{k-1}
\end{pmatrix}
\]

and therefore the Jacobian \(A_k\) is

\[
A_k = \begin{pmatrix}
1 & -\frac{1}{2} \Delta t & 0 & 0 & 0 & 0 \\
0 & 1 - \theta \Delta t & \Delta t & -v_{k-1} \Delta t & 0 & 0 \\
0 & 0 & 1 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & 0 \\
0 & 0 & 0 & 0 & 0 & 1
\end{pmatrix}
\]

\(^{16}\)In reality we should write the estimation parameters \(\omega_k\), \(\theta_k\), \(\xi_k\) and \(\rho_k\). However we drop the indexes for simplifying the notations.
and

$$W_k = \begin{pmatrix} \sqrt{v_{k-1}} \Delta t & 0 \\ 0 & \xi \sqrt{v_{k-1}} \Delta t \end{pmatrix}$$

having the measurement $z_k = \ln S_k$ we can write

$$H_k = \begin{pmatrix} 1 & 0 & 0 & 0 & 0 \end{pmatrix}$$

and $U_k = 0$.

We could however introduce a measurement noise $R$ corresponding to the intra-day stock price movements and the bid-ask spread, in which case we would have $z_k = \ln S_k + R \epsilon_k$ where $\epsilon_k$ represents a sequence of uncorrelated standard Normal random variables. This means that $R_k = R$ and $U_k = 1$. We can then tune the value of $R$ in order to get more stable results.

**Example 2** The same exact methodology could be used in the GARCH framework. We define the state variable $x^k_t = (\ln S_k, v_k, \omega_0, \alpha, \beta, c)$ and taking for observation the logarithm of the actual stock price $S_k$.

The system could be written as

$$x_k = f(x_{k-1}, w_{k-1})$$

with $w_k = B_k$ a one-dimensional source of noise with a variance $Q_k = 1$ and

$$f(x_{k-1}, w_{k-1}) = \begin{pmatrix} \ln S_{k-1} + (\mu_S - \frac{1}{2} v_{k-1}) + \sqrt{v_{k-1}} B_{k-1} \\ \omega_0 + \beta v_{k-1} + \alpha (B_{k-1} - c \sqrt{v_{k-1}})^2 \\ \omega_0 \\ \alpha \\ \beta \\ c \end{pmatrix}$$

and the Jacobian

$$A_k = \begin{pmatrix} 1 & -\frac{1}{2} & 0 & 0 & 0 & 0 \\ 0 & \beta + \alpha c^2 & 1 & c^2 v_{k-1} & v_{k-1} & 2 \alpha c v_{k-1} \\ 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 \end{pmatrix}$$
and

\[ W_k = \begin{pmatrix} \sqrt{v_{k-1}} \\ -2\alpha c \sqrt{v_{k-1}} \\ 0 \\ 0 \\ 0 \\ 0 \end{pmatrix} \]

The observation \( z_k \) will be

\[ z_k = h(x_k) = \ln(S_k) \]

exactly as in the previous example. The rest of the algorithm would therefore be identical to the one included in Example 1.

**Example 3** In the above example we included all the variables in the system process and we observed part of the system. It is also possible to separate the measurement and the system variables as follows.

Taking a general discrete Stochastic Volatility process as

\[ \ln S_k = \ln S_{k-1} + (\mu_S - \frac{1}{2}v_k)\Delta t + \sqrt{v_k}\Delta t B_k \]

\[ v_k = v_{k-1} + b(v_{k-1})\Delta t + a(v_{k-1})\sqrt{\Delta t}Z_k \]

with \( B_k \) and \( Z_k \) two Normal random sequences with a mean of zero and variance one, with a correlation equal to \( \rho \).

Posing \( y_k = \sqrt{v_k}Z_k \) and performing the usual Cholesky factorization \( B_k = \rho Z_k + \sqrt{1 - \rho^2}X_k \) where \( Z_k \) and \( X_k \) are uncorrelated, we can now take the case of a Square-Root process and write

\[ x_k = \begin{pmatrix} v_k \\ y_k \\ \omega \\ \theta \\ \xi \\ \rho \end{pmatrix} \]

and \( x_k = f(x_{k-1}, Z_k) \) with

\[ f(x_{k-1}, Z_k) = \begin{pmatrix} v_{k-1} + (\omega - \theta v_{k-1})\Delta t + \xi \sqrt{v_{k-1}}\sqrt{\Delta t}Z_k \\ (v_{k-1} + (\omega - \theta v_{k-1})\Delta t + \xi \sqrt{v_{k-1}}\sqrt{\Delta t}Z_k)^2 Z_k \\ \omega + QZ_k \\ \theta + QZ_k \\ \xi + QZ_k \\ \rho + QZ_k \end{pmatrix} \]

\(^{17}\)Note that the indexing here is slightly different from the previous examples.
which provides us with the Jacobian

\[ A_k = \begin{pmatrix}
1 - \theta \Delta t & \Delta t & -v_{k-1} \Delta t & 0 & 0 \\
0 & 0 & 0 & 0 & 0 \\
0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 \\
0 & 0 & 0 & 0 & 1 \\
\end{pmatrix} \]

and

\[ W_k = \begin{pmatrix}
\xi \sqrt{v_{k-1}} \sqrt{\Delta t} \\
(v_{k-1} + (\omega - \theta v_{k-1}) \Delta t)^{\frac{3}{2}} \\
Q \\
Q \\
Q \\
Q \\
\end{pmatrix} \]

The measurement equation is

\[ z_k = \ln\left(\frac{S_k}{S_{k-1}}\right) = (\mu_S - \frac{1}{2} v_k) \Delta t + \rho \sqrt{\Delta t} y_k + \sqrt{1 - \rho^2} \sqrt{\Delta t} \sqrt{1 - \rho^2} \sqrt{\Delta t} X_k \]

and therefore

\[ H_k = \begin{pmatrix}
-\frac{1}{2} \Delta t & \rho \sqrt{\Delta t} & 0 & 0 & 0 \\
\end{pmatrix} \]

with \( u_k = X_k \) and \( U_k = \sqrt{1 - \rho^2} \sqrt{\Delta t} \) which completes our set of equations. Again we could tune the system noise \( Q \) in order to obtain more stable results.
Figure 2.7: EKF Estimation (Example 1) for the drift parameter $\omega$
SPX Index Daily Close Prices were used over five years from 19961001 to 20010928. The convergence is fairly good.
Figure 2.8: EKF Estimation (Example 1) for the drift parameter $\theta$

SPX Index Daily Close Prices were used over five years from 19961001 to 20010928. The convergence is fairly good.
Figure 2.9: EKF Estimation (Example 1) for the volatility-of-volatility parameter $\xi$.
SPX Index Daily Close Prices were used over five years from 19961001 to 20010928. The convergence is rather poor. We shall explain this via the concept of observability.
Figure 2.10: EKF Estimation (Example 1) for the correlation parameter $\rho$

SPX Index Daily Close Prices were used over five years from 19961001 to 20010928. The convergence is rather poor. We shall explain this via the concept of observability.
Observability

From the above tests it seems that the EKF provides us with a non-robust calibration methodology. Indeed the results are very sensitive to the choice of system noise $Q$ and observation noise $R$. We chose for this case $Q = 10^{-3}$ and $R \approx 0$.

This brings to attention the issue of Observability. A non-linear system with a state vector $x_k$ of dimension $n$ is observable if

$$O = \begin{pmatrix} H \\ HA \\ HA^2 \\ \vdots \\ HA^{n-1} \end{pmatrix}$$

has a full rank of $n$. For an explanation refer to Reif et al. [205].

It is fairly easy to see that among the above examples, the first and third one (corresponding to the Stochastic Volatility formulation) have for the observation matrix $O$ a rank of four and therefore are not observable. This explains why they do not converge well and are so sensitive to the tuning parameters $Q$ and $R$.

This means that the choices of the state variables for the Examples 1 and 3 were rather poor. One reason is that in our state space choice, we considered

$$z_k = h(v_{k-1}, ...)$$

and

$$x_k = (..., v_k, ...) = f(x_{k-1}, ...)$$

which implies that

$$\frac{\partial h}{\partial v_k} = 0$$

We shall see how to correct this in the next section by choosing a more appropriate state space representation.
The One-Dimensional State within the Joint Filter

Considering the state equation

\[ v_k = v_{k-1} + (\omega - \theta v_{k-1}) \Delta t + \xi \sqrt{v_{k-1}} \sqrt{\Delta t} Z_{k-1} - \rho \xi \ln S_{k-1} + (\mu_S - \frac{1}{2} v_{k-1}) \Delta t + \sqrt{v_{k-1}} \sqrt{\Delta t} B_{k-1} - \ln S_k \]

posing for every \( k \)

\[ \tilde{Z}_k = \frac{1}{\sqrt{1 - \rho^2}} (Z_k - \rho B_k) \]

we will have as expected \( \tilde{Z}_k \) uncorrelated with \( B_k \). Therefore considering the augmented state

\[ x_k = \begin{pmatrix} v_k \\ \omega \\ \theta \\ \xi \\ \rho \end{pmatrix} \]

we will have the state transition equation

\[ f(x_{k-1}, \tilde{Z}_{k-1}) = \begin{pmatrix} v_{k-1} + [(\omega - \rho \xi) v_{k-1}] \Delta t + \rho \xi \ln \left( \frac{S_k}{S_{k-1}} \right) + \xi \sqrt{1 - \rho^2} \sqrt{v_{k-1}} \sqrt{\Delta t} \tilde{Z}_{k-1} \\ \omega \\ \theta \\ \xi \\ \rho \end{pmatrix} \]

and the measurement equation would be

\[ z_k = \ln S_{k+1} = \ln S_k + (\mu_S - \frac{1}{2} v_k) \Delta t + \sqrt{v_k} \sqrt{\Delta t} B_k \]

The corresponding EKF Jacobians for this system are

\[ A_k = \begin{pmatrix} 1 - (\theta - \frac{1}{2} \rho \xi) \Delta t & \Delta t & -v_{k-1} \Delta t & \rho \left( \ln \left( \frac{S_k}{S_{k-1}} \right) - (\mu_S - \frac{1}{2} v_{k-1}) \Delta t \right) & \xi \left( \ln \left( \frac{S_k}{S_{k-1}} \right) - (\mu_S - \frac{1}{2} v_{k-1}) \Delta t \right) \\ 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 & 1 \end{pmatrix} \]

\[ W_k = \begin{pmatrix} \xi \sqrt{1 - \rho^2} \sqrt{v_{k-1}} \sqrt{\Delta t} \\ 0 \\ 0 \\ 0 \\ 0 \end{pmatrix} \]
\[ H_k = \begin{pmatrix} -\frac{1}{2} \Delta t & 0 & 0 & 0 \end{pmatrix} \]
\[ U_k = \sqrt{v_k} \sqrt{\Delta t} \]

It is easy to check that this system is observable since the observation matrix \( O_k \) is of full rank. This shows that our state space choice is better than the previous ones.

The UKF would be implemented in a similar fashion from the transition and observation equations above. Again, for the UKF we would *not* need to compute any Jacobians.

An important issue to consider is that of *Tuning*. We could add extra-noise to the observation and hence lower the weight associated with the observations. In which case after choosing a tuning parameter \( R \) we would write
\[ U_k = \begin{pmatrix} \sqrt{v_k} \sqrt{\Delta t} & R \end{pmatrix} \]

and
\[ U_k U_k^T = v_k \Delta t + R^2 \]

The choice of the initial conditions and the tuning parameters could make the algorithm fail or succeed. It therefore seems that there is little robustness in this procedure.

We consider the example of 5000 artificial data points artificially produced via a Heston stochastic volatility process with a parameter-set
\[ \Psi^* = (\omega = 0.10, \theta = 10.0, \xi = 0.03, \rho = -0.50) \]

with a given \( \mu_S = 0.025 \). We then choose a tuning parameter \( R = 0.10 \) and take a reasonable guess for the initial-conditions
\[ \Psi_0 = (\omega_0 = 0.15, \theta_0 = 10.0, \xi_0 = 0.02, \rho_0 = -0.51) \]

and apply the Joint Filter. The results are displayed in figures 2.11 to 2.14. As we can see, the convergence for \( \omega \) and \( \theta \) are better than the two others. We shall see later why this is.

Allowing a *burn in* period of 1000 points, we can calculate the mean (and the standard-deviation) of the generated parameters after the simulation 1000.
Figure 2.11: Joint EKF Estimation for the parameter $\omega$
Prices were simulated with $\Psi^* = (0.10, 1.0, 0.03, -0.50)$. The convergence remains mediocre. We shall explain this in the following section.
Figure 2.12: Joint EKF Estimation for the parameter $\theta$
Prices were simulated with $\Psi^* = (0.10, 10.0, 0.03, -0.50)$. The convergence remains mediocre. We shall explain this in the following section.
Figure 2.13: Joint EKF Estimation for the parameter $\xi$

Prices were simulated with $\Psi^* = (0.10, 10.0, 0.03, -0.50)$. The convergence remains mediocre. We shall explain this in the following section.
Prices were simulated with $\Psi^* = (0.10, 10.0, 0.03, -0.50)$. The convergence remains mediocre. We shall explain this in the following section.
Joint Filters and Time Interval

One difficulty with the application of the Joint Filter (JF) to the Stochastic Volatility problem, is the following: Unless we are dealing with a longer time-interval such as $\Delta t = 1$, the observation error $\sqrt{v_k} \sqrt{\Delta t} B_k$ is too large compared to the sensitivity of the filter with respect to the state through $-0.5 v_k \Delta t$. Indeed for a $\Delta t = 1/252$ we have

$$\Delta t = o(\sqrt{\Delta t})$$

A simple Monte-Carlo test will allow us to verify this: We simulate a Heston model on the one hand, and a modified model where we multiply both Brownian Motions by a factor $\Delta t$. This will make the errors smaller by a factor of 252 for the daily case. We call this model the Modified Model.

After generating 5000 data points with a parameter set $(\omega^* = 0.10, \theta^* = 10.0, \xi^* = 0.03, \rho^* = -0.50)$ and a drift $\mu_S = 0.025$, we suppose we know all parameters except $\omega$.

We then apply the JKF to find the estimate $\hat{\omega}$. We can observe in figure 2.15 that the filter diverges when applied to the Heston model but converges fast when applied to the Modified model.

However in reality we have no control over the observation error, which is precisely the volatility!

In a way, this brings up a more fundamental issue regarding the stochastic volatility Estimation problem: By definition, volatility represents the noise of the stock process. Indeed if we had taken the spot price $S_k$ as the observation and the variance $v_k$ as the state, we would have

$$S_k = S_{k-1} + S_{k-1} \mu_S \Delta t + S_{k-1} \sqrt{v_k} \sqrt{\Delta t} B_k$$

we would then have an observation function gradient $H = 0$ and the system would be unobservable!

It is precisely because we use a Taylor second order expansion

$$\ln(1 + R) \approx R - \frac{1}{2} R^2$$

that we obtain access to $v_k$ through the observation function. However the error remains dominant as the first order of the expansion.

Some [130] have tried

$$\ln \left( \ln^2 \left( \frac{S_k}{S_{k-1}} \right) \right) \approx \ln(v_k) + \ln(\Delta t) + \ln(B_k^2)$$

and

$$\ln(B_k^2) \sim -1.27 + \frac{\pi}{\sqrt{2}} \mathcal{N}(0, 1)$$

\[130\] Hereafter $x_h = o(y_h)$ means $x_h/y_h \to 0$ as $h \to 0$, or more intuitively, $x_h$ is much smaller than $y_h$ for a tiny $h$. 

\[131\]
but the latter approximation may or may not be valid depending on the problem under study.
Figure 2.15: Joint EKF Estimation for the parameter $\omega$ applied to the Heston model as well as to a Modified model where the noise is reduced by a factor 252. As we can see, the convergence for the modified model is improved dramatically. This justifies our comments on large observation error.
2.3.7 Parameter Estimation via MLE

As previously stated, one of the principal methods of estimation under the classical framework is the Maximization of the Likelihood. Indeed this estimation method has many desirable asymptotic properties.

Therefore instead of using the filters alone, we could separate the parameter-set $\Psi = (\omega, \theta, \xi, \rho)$ from the state vector $(\ln S_k, v_k)$ and use the Kalman Filter for state filtering within each MLE iteration\(^\dagger\) and estimate the parameters iteratively.

**An Illustration**

Let us first consider the case of the previous illustration

$$\xi_k = \xi_{k-1} + \pi + 0.10 w_k$$

and

$$z_k = \xi_k + 0.10 u_k$$

where $\pi \approx 3.14159$ and $w_k, u_k$ are independent Gaussian random variables.

Here we take

$$x_k = \xi_k$$

and

$$A_k = H_k = 1$$

$$W_k = U_k = 0.1$$

The maximization of the Gaussian Likelihood with respect to the parameter $\pi$ is equivalent to minimizing

$$L_{1:N} = \sum_{k=1}^{N} \left[ \ln(F_k) + \frac{z_k^2}{F_k} \right]$$

with residuals

$$\tilde{z}_k = z_k - \hat{x}_k = z_k - \hat{\xi}_k$$

and

$$F_k = P_{z_k z_k} = H_k P_{\xi_k} H_k^T + U_k R_k U_k^T$$

Note that we used *scalar* notations here and in vectorial notations we would have

$$L_{1:N} = \sum_{k=1}^{N} \left[ \ln(|F_k|) + \tilde{z}_k^T F_k^{-1} \tilde{z}_k \right]$$

\(^\dagger\)To be more accurate, since the noise process is *conditionally* Gaussian, we are dealing with a Quasi-Maximum-Likelihood (QML) Estimation. More detail could be found for instance in Gourieroux [124].
where $\vert \mathbf{F}_k \vert$ is the determinant of $\mathbf{F}_k$.

We use the scalar notations for simplicity and also because in the Stochastic Volatility problem, we usually deal with one-dimensional observations (namely the Stock Price).

The minimization via a Direction Set (Powell) method over 500 artificially generated observation points will provide

$$\hat{\pi} = 3.145953$$

very quickly.

**Stochastic Volatility Examples**

For our previous first example the system state vector now becomes

$$\mathbf{x}_k = \begin{pmatrix} \ln S_k \\ v_k \end{pmatrix}$$

which means the dimension of our state is now two, and

$$\mathbf{x}_k = \mathbf{f}(\mathbf{x}_{k-1}, \mathbf{w}_{k-1}) = \begin{pmatrix} \ln S_{k-1} + (\mu S - \frac{1}{2} v_{k-1}) \Delta t + \sqrt{v_{k-1}} \sqrt{\Delta t} B_{k-1} \\ v_{k-1} + (\omega - \theta v_{k-1}) \Delta t + \xi \sqrt{v_{k-1}} \sqrt{\Delta t} Z_{k-1} \end{pmatrix}$$

The system noise is still

$$\mathbf{w}_k = \begin{pmatrix} B_k \\ Z_k \end{pmatrix}$$

and the corresponding covariance matrix is

$$\mathbf{Q}_k = \begin{pmatrix} 1 & \rho \\ \rho & 1 \end{pmatrix}$$

We have the measurement $\mathbf{z}_k = \ln S_k$ and therefore we can write

$$\mathbf{H}_k = \begin{pmatrix} 1 & 0 \end{pmatrix}$$

Now for a given set of parameters $(\omega, \theta, \xi, \rho)$ we can filter this system with the EKF (or the UKF) using

$$\mathbf{A}_k = \begin{pmatrix} 1 & -\frac{1}{2} \Delta t \\ 0 & 1 - \theta \Delta t \end{pmatrix}$$

and

$$\mathbf{W}_k = \begin{pmatrix} \sqrt{v_{k-1}} \sqrt{\Delta t} & 0 \\ 0 & \xi \sqrt{v_{k-1}} \sqrt{\Delta t} \end{pmatrix}$$

Note that the observation matrix is

$$\mathbf{O}_k = \begin{pmatrix} 1 & 0 \\ 1 & -\frac{1}{2} \Delta t \end{pmatrix}$$
which is of full rank. Our system is therefore observable.

After filtering for this set of parameters, we calculate the sum to be minimized

\[ \phi(\omega, \theta, \xi, \rho) = \sum_{k=1}^{N} \left[ \ln(F_k) + \frac{\tilde{z}_k^2}{F_k} \right] \]

with

\[ \tilde{z}_k = z_k - h(\hat{x}_k^{-}, 0) \]

and

\[ F_k = H_k P_k^{-} H_k^T + U_k R_k U_k^T \]

The minimization could once again be done via a Direction Set (Powell) method as described previously. This will avoid a calculation of the gradient \( \nabla \phi \).

It is interesting to observe (cf. figures 2.16 and 2.17) that the results of the EKF and UKF are very close and the filter errors are comparable. However the estimated parameter set \( \Psi = (\omega, \theta, \xi, \rho) \) can have a different set of values depending on which filter is actually used.\(^{20}\)

Hence the question, how sensitive are these filters to \( \Psi \)? In order to answer this question we can run an estimation for EKF and use the estimated parameters in UKF and observe how good a fit we obtain. The results show that this sensitivity is fairly low. Again, this might be due to the relative insensitivity of the filters to the parameter-set \( \Psi \) or the non-uniqueness of the optimal parameter-set. As we will see, the answer to this question also depends on the sample size.

**Optimization-Constraints for the Square-Root Model**

In terms of the optimization constraints, in addition to the usual

\[ \omega \geq 0 \]  \hspace{1cm} (2.12)

\[ \theta \geq 0 \]

\[ \xi \geq 0 \]

\[ -1 \leq \rho \leq 1 \]

we need to make sure that the value of the variance remains positive, i.e.

\[ v_k + (\omega - \theta v_k) \Delta t + \xi \sqrt{v_k} \sqrt{\Delta t} Z_k \geq 0 \]

for any \( v_k \geq 0 \) and any Gaussian random value \( Z_k \).

For a Gaussian random variable \( Z_k \) and any positive real number \( Z^* \), we \(^{20}\)Note however that the values of the resulting long-term volatilities \( \sqrt{v_k} \) are rather close.
can write $Z_k \geq -Z^*$ with a probability $P^*$. For instance if $Z^* = 4$ then $P^* = 0.999968$. Therefore, fixing a choice of $Z^*$, it is almost always enough for us to have

$$v_k + (\omega - \theta v_k) \Delta t - \xi \sqrt{v_k} \sqrt{\Delta t} Z^* \geq 0$$

for any $v_k \geq 0$.

Considering the function $f(x) = x + (\omega - \theta x) \Delta t - \xi \sqrt{x} \sqrt{\Delta t} Z^*$ it is fairly easy to see that $f(0) = \omega \Delta t \geq 0$ by assumption, and for $x$ very large $f(x) \approx (1 - \theta \Delta t)x$ which is positive if

$$\theta \leq \frac{1}{\Delta t} \quad (2.13)$$

This is most of the time realized for a small $\Delta t$ such as ours.

Now $f(x)$ being a continuous function and having positive values at zero and infinity it would be sufficient to make sure that its one minimum on $[0, +\infty[$ is also positive. A simple derivative computation shows that $x_{\text{min}} = \frac{\xi^2 \Delta t Z^*}{4(1 - \theta \Delta t)^2}$ and therefore the positivity is realized if$^{21}$

$$\xi \leq \frac{2}{Z^*} \sqrt{\omega(1 - \theta \Delta t)} \quad (2.14)$$

which completes our set of constraints.

---

$^{21}$Naturally we suppose $\Delta t > 0$
Figure 2.16: SPX Historic Data (1996-2001) is filtered via EKF and UKF. The results are very close however the estimated parameters $\hat{\Psi} = (\hat{\omega}, \hat{\theta}, \hat{\xi}, \hat{\rho})$ differ. Indeed we find $\hat{\omega} = 0.073028$, $\hat{\theta} = 1.644488$, $\hat{\xi} = 0.190461$, $\hat{\rho} = -1.000000$) for the EKF and $(\hat{\omega} = 0.540715, \hat{\theta} = 13.013577, \hat{\xi} = 0.437523, \hat{\rho} = -1.000000)$ for the UKF. This might be due to the relative insensitivity of the filters to the parameter-set $\Psi$ or the non-uniqueness of the optimal parameter-set. We shall explain this low sensitivity in more detail.
Figure 2.17: EKF and UKF absolute filtering-errors for the same time-series. As we can see there is no clear superiority of one algorithm over the other.
An Alternative Implementation

We could also perform the same estimation but based on our previous third example. Again we have

\[
\ln S_k = \ln S_{k-1} + (\mu_S - \frac{1}{2} v_k) \Delta t + \sqrt{v_k \Delta t} B_k
\]

\[
v_k = v_{k-1} + (\omega - \theta v_{k-1}) \Delta t + \xi \sqrt{v_{k-1} \Delta t} Z_k
\]

with \( B_k \) and \( Z_k \) two Normal random sequences with a mean of zero, a variance of one, and a correlation equal to \( \rho \).

However since for a Kalman Filter the process noise and the measurement noise must be uncorrelated, we introduce

\[
y_k = \sqrt{v_k} Z_k
\]

and performing the usual Cholesky factorization \( B_k = \rho Z_k + \sqrt{1 - \rho^2} X_k \) where \( Z_k \) and \( X_k \) are uncorrelated, we can write

\[
x_k = \begin{pmatrix} v_k \\ y_k \end{pmatrix}
\]

and \( x_k = f(x_{k-1}, Z_k) \) with

\[
f(x_{k-1}, Z_k) = \begin{pmatrix} v_{k-1} + (\omega - \theta v_{k-1}) \Delta t + \xi \sqrt{v_{k-1} \Delta t} Z_k \\ v_{k-1} + (\omega - \theta v_{k-1}) \Delta t + \xi \sqrt{v_{k-1} \Delta t} Z_k \end{pmatrix}
\]

which provides us with the Jacobian

\[
A_k = \begin{pmatrix} 1 - \theta \Delta t & 0 \\ 0 & 0 \end{pmatrix}
\]

and

\[
W_k = \begin{pmatrix} \xi \sqrt{v_{k-1} \Delta t} \\ \sqrt{v_{k-1} \Delta t} \end{pmatrix}
\]

The measurement equation is

\[
z_k = \ln S_k = \ln S_{k-1} + (\mu_S - \frac{1}{2} v_k) \Delta t + \rho \sqrt{\Delta t} y_k + \sqrt{1 - \rho^2} \sqrt{v_k \Delta t} X_k
\]

and therefore

\[
H_k = \begin{pmatrix} -\frac{1}{2} \Delta t & \rho \sqrt{\Delta t} \\ -\frac{1}{2} \Delta t & \rho \sqrt{\Delta t} \end{pmatrix}
\]

with \( u_k = X_k \) and \( U_k = \sqrt{1 - \rho^2} \sqrt{v_k \Delta t} \) which completes our set of equations.

Note that the observation matrix is

\[
O_k = \begin{pmatrix} -\frac{1}{2} \Delta t & \rho \sqrt{\Delta t} \\ -\frac{1}{2} \Delta t (1 - \theta \Delta t) & 0 \end{pmatrix}
\]

which is of full rank. Our system is therefore observable.
The One-Dimensional State

Finally, a simpler way of writing the state-space system, which will be our method of choice hereafter, would be to subtract from both sides of the state equation $x_k = f(x_{k-1}, w_{k-1})$, a multiple of the quantity $h(x_{k-1}, u_{k-1}) - z_{k-1}$ which is equal to zero. This would allow us to eliminate the correlation between the system and the measurement noises.

Indeed, if the system equation is

$$
\ln S_k = \ln S_{k-1} + (\mu_S - \frac{1}{2}v_{k-1}) \Delta t + \sqrt{v_{k-1}} \sqrt{\Delta t}B_{k-1}
$$

$$
v_k = v_{k-1} + (\omega - \theta v_{k-1}) \Delta t + \xi \sqrt{v_{k-1}} \sqrt{\Delta t}Z_{k-1}
$$

writing

$$
v_k = v_{k-1} + (\omega - \theta v_{k-1}) \Delta t + \xi \sqrt{v_{k-1}} \sqrt{\Delta t}Z_{k-1} - \rho \xi \ln S_k
$$

posing for every $k$

$$
\hat{Z}_k = \frac{1}{\sqrt{1 - \rho^2}} (Z_k - \rho B_k)
$$

we will have as expected $\hat{Z}_k$ uncorrelated with $B_k$ and

$$
x_k = v_k = v_{k-1} + [(\omega - \rho \xi \mu_S) - (\theta - \frac{1}{2} \rho \xi) v_{k-1}] \Delta t + \rho \xi \ln(\frac{S_k}{S_{k-1}}) + \xi \sqrt{1 - \rho^2} \sqrt{v_{k-1}} \sqrt{\Delta t} \hat{Z}_{k-1}
$$

and the measurement equation would be

$$
z_k = \ln S_{k+1} = \ln S_k + (\mu_S - \frac{1}{2}v_k) \Delta t + \sqrt{v_k} \sqrt{\Delta t}B_k
$$

With this system everything becomes one-dimensional and the computations become much faster both for the EKF and UKF.

For the EKF we will have

$$
A_k = 1 - (\theta - \frac{1}{2} \rho \xi) \Delta t
$$

and

$$
W_k = \xi \sqrt{1 - \rho^2} \sqrt{v_{k-1}} \sqrt{\Delta t}
$$

as well as

$$
H_k = -\frac{1}{2} \Delta t
$$

and

$$
U_k = \sqrt{v_k} \sqrt{\Delta t}
$$
Again, for the MLE we will try to minimize

$$\phi(\omega, \theta, \xi, \rho) = \sum_{k=1}^{N} \left[ \ln(F_k) + \frac{\tilde{z}_k^2}{F_k} \right]$$

with residuals

$$\tilde{z}_k = z_k - h(\hat{x}_k^-, 0)$$

and

$$F_k = H_k P_k^- H_k^\dagger + U_k U_k^\dagger$$

The same time update and measurement update will be used with the UKF. The ML Estimator can be used as usual.

**Other stochastic volatility models**

It is easy to generalize the above state space model to other stochastic volatility approaches. Indeed we could replace the Heston equation with

$$v_k = v_{k-1} + (\omega - \theta v_{k-1}) \Delta t + \xi v_{k-1}^p \sqrt{\Delta t} Z_k$$

where $p = 1/2$ would naturally correspond to the Heston model, $p = 1$ to the GARCH diffusion-limit model, and $p = 3/2$ to the $\frac{3}{2}$ model described in [177]. This idea will be developed further in the chapter.
2.3.8 Diagnostics

After having estimated the parameter-set $\Psi$ we should test for model misspecification. Two important questions are

1. Do the normalized residuals $(z_k - \hat{z}_k)/F_k$ follow a standard normal $N(0, 1)$ law?

2. Do these residuals have zero auto-correlation?

**Chi-Square Test**

The first question could be answered by performing a Chi-Square test. We take a number $N_B$ of intervals or “bins” bounded by the points $x_0, x_1, ..., x_J$. We then count the number of observations $N_j$ within each bin $[x_j, x_{j+1}]$ for $j$ between zero and $N_B - 1$. We then compare these numbers with the theoretical numbers implied by the normal distribution $n_j = [\Phi(x_{j+1}) - \Phi(x_j)]N$ with $\Phi$ the cumulative normal function and $N$ the total number of observations. The sum

$$\sum_{j=0}^{N_B-1} \frac{(N_j - n_j)^2}{n_j}$$

asymptotically follows a $\chi^2$ law with degrees of freedom $\nu$ equal to $N_B - 1$.

**Box-Ljung Test**

The second question could be answered with a Box-Ljung test. We should first calculate a number of auto-correlations

$$r_k = \frac{\sum_{i=1}^{N-k}(\tilde{z}_i - \bar{\tilde{z}})(\tilde{z}_{i+k} - \bar{\tilde{z}})}{\sum_{i=1}^{N}(\tilde{z}_i - \bar{\tilde{z}})^2}$$

for $k$ between one and a prespecified integer $K$. Once again $\tilde{z}_i = z_i - \hat{z}_i$ and $\bar{\tilde{z}}$ corresponds to their mean.

We then consider the sum

$$N(N + 2) \sum_{k=1}^{K} \frac{r_k^2}{N - k}$$

which asymptotically follows a $\chi^2_\nu$ law with degrees of freedom $\nu$ equal to $K - p$ where $p = 4$ is the number of parameters we estimated.
Test Results

In the previously studied SPX examples we had \( N = 1256 \). For the Normality test we choose \( N_B = 21 \) and for the Box-Ljung test we take \( K = 24 \), in both cases we will have to compare the outputs to the critical threshold \( \chi^2_{20} \) which for a confidence of 0.95 is around 31.5.

For the (one-dimensional) EKF we obtain 27.738862 for the Normality test and 0.007889 for the Box-Ljung test. For the (one-dimensional) UKF we obtain 22.657545 for the Normality test and again 0.016053 for the Box-Ljung test.

This means that there is very little auto-correlation in our system noise. Also, it seems reasonable to model the noise as approximately normally distributed.

The Chi-Square test proves that the Normality assumption is plausible and the Kalman Filter can indeed be used.

A visual confirmation of this could be achieved by plotting the corresponding histograms. As we can see in figure 2.18, there are no excessively “fat tails”, however the central value at zero is higher than the normal distribution.
Figure 2.18: Histogram for filtered data via EKF versus the Normal distribution. The residuals are fairly normal.
Variogram

Similarly to Fouque et al [104] we can use a variogram to visualize the volatility behavior of the model. As Galli [110] mentions, the main reasons to use variograms instead of covariance or correlogram are: variograms do not need to estimate the mean, and they are interpretable under wider conditions than covariances or correlograms.

The expression for the variogram of a process $I_t$ is

$$
\gamma_I(h) = \frac{1}{2} \mathbb{E}[(I(t + h) - I(t))^2] \approx \frac{1}{2N_{h_i}} \sum_{t=0}^{N_{h_i}} (I(t + h_i) - I(t))^2
$$

where $N_{h_i}$ is the total number of points such that $I(t + h_i)$ exists.

For instance for a sequence of Independent Gaussian Random-Variables we should have

$$
\gamma_I(h) = \frac{1}{2} \mathbb{E}[I^2(t + h)] + \frac{1}{2} \mathbb{E}[I^2(t)] - \mathbb{E}[I(t)I(t + h)] = \frac{1}{2} + \frac{1}{2} - 0 = 1
$$

In our case, the process $I_t$ could be defined for instance as

$$
I_t = \frac{z_t - \hat{z}_t}{\sqrt{F_t}}
$$

which should correspond to a sequence of Independent Gaussian Random-Variables.

As we can see in figure 2.19, the variogram is consistent with the Gaussian assumption which reconfirms what we observed from the histograms.

Another way of expressing the same idea is to build a Brownian Motion from the above sequence. Calling the Independent Gaussian Random-Variables $(B_k)$, we can write

$$
I_n = \sqrt{\Delta t} \sum_{k=0}^{n} B_k
$$

and plot the variogram for the Brownian Motion $I_n$.

For a Brownian Motion, it is easy to see that the variogram should be linear

$$
\gamma_I(h) = \frac{1}{2} (t + h) + \frac{1}{2} t - t = \frac{1}{2} h
$$

This could indeed be seen in figure 2.20.
Figure 2.19: Variograms for filtered data via EKF and UKF. The input corresponds to a sequence of Independent Gaussian Random-Variables. As we can see, the variograms are close to one.
Figure 2.20: Variograms for filtered data via EKF and UKF. The input corresponds to a Brownian Motion. As we can see, the Variograms are close to $x/2$. 
2.3.9 Particle Filtering

A different approach to filtering and parameter estimation has recently become popular [79], [122], [171]. In this approach, we use Monte-Carlo Simulations instead of Gaussian approximations for \((x_k|z_k)\) as the Kalman or Kushner Filters do. This will precisely allow us to deal with fundamentally Non-Gaussian situations.\(^{22}\)

Underlying Theory

The idea is based on the Importance Sampling technique:

We can calculate an expected value

\[
\mathbb{E}[f(x_{0:k})] = \int f(x_{0:k}) p(x_{0:k}|z_{1:k}) dx_{0:k}
\]

by using a known and simple proposal distribution \(q()\).

More precisely, it is possible to write

\[
\mathbb{E}[f(x_{0:k})] = \int f(x_{0:k}) \frac{p(x_{0:k}|z_{1:k})}{q(x_{0:k}|z_{1:k})} q(x_{0:k}|z_{1:k}) dx_{0:k}
\]

which could be also written as

\[
\mathbb{E}[f(x_{0:k})] = \int f(x_{0:k}) \frac{w_k(x_{0:k})}{p(z_{1:k})} q(x_{0:k}|z_{1:k}) dx_{0:k}
\]

with

\[
w_k(x_{0:k}) = \frac{p(z_{1:k}|x_{0:k}) p(x_{0:k})}{q(x_{0:k}|z_{1:k})}
\]

defined as the filtering non-normalized weight as step \(k\).

Proof:

Indeed, we can write

\[
\frac{p(x_{0:k}|z_{1:k})}{q(x_{0:k}|z_{1:k})} = \frac{p(z_{1:k}|x_{0:k}) p(x_{0:k})}{p(z_{1:k}) q(x_{0:k}|z_{1:k})}
\]

\[
= \frac{w_k(x_{0:k})}{p(z_{1:k})}
\]

as we claimed. (QED)

We therefore have

\[
\mathbb{E}[f(x_{0:k})] = \mathbb{E}_q[\frac{w_k(x_{0:k}) f(x_{0:k})}{w_k(x_{0:k})}] = \mathbb{E}_q[\tilde{w}_k(x_{0:k}) f(x_{0:k})]
\]

\(^{22}\)An existing (but less effective) alternative to the Particle Filtering method is the Grid-based Approximation as the one suggested by Kitagawa [170], [108]. The main advantage of the Particle Filter is that it will make the grid focus adaptively on the state-space regions with higher relevance.
with

\[ \tilde{w}_k(x_{0:k}) = \frac{w_k(x_{0:k})}{\mathbb{E}_q[w_k(x_{0:k})]} \]

defined as the filtering \textit{normalized weight} as step \( k \).

\textbf{Proof:}

Indeed

\[
\mathbb{E}[f(x_{0:k})] = \frac{1}{p(z_{1:k})} \int f(x_{0:k}) w_k(x_{0:k}) q(x_{0:k}|z_{1:k}) dx_{0:k}
\]

\[
= \int \frac{f(x_{0:k}) w_k(x_{0:k}) q(x_{0:k}|z_{1:k})}{p(z_{1:k}|x_{0:k}) p(x_{0:k}) q(x_{0:k}|z_{1:k})} dx_{0:k}
\]

\[
= \frac{\int f(x_{0:k}) w_k(x_{0:k}) q(x_{0:k}|z_{1:k}) dx_{0:k}}{\int w_k(x_{0:k}) q(x_{0:k}|z_{1:k}) dx_{0:k}}
\]

which is the ratio of the expectations as earlier stated. \((QED)\)

Using Monte-Carlo sampling from the distribution \( q(x_{0:k}|z_{1:k}) \) we can write in the discrete framework:

\[
\mathbb{E}[f(x_{0:k})] \approx \frac{1}{N_{\text{sims}}} \sum_{i=1}^{N_{\text{sims}}} \tilde{w}_k(x_{0:k}^{(i)}) f(x_{0:k}^{(i)})
\]

with again

\[
\tilde{w}_k(x_{0:k}^{(i)}) = \frac{w_k(x_{0:k}^{(i)})}{\sum_{j=1}^{N_{\text{sims}}} w_k(x_{0:k}^{(j)})}
\]

Now supposing that our proposal distribution \( q() \) satisfies the Markov property, it can be shown that \( w_k \) verifies the recursive identity

\[
w_k^{(i)} = w_{k-1}^{(i)} \frac{p(z_k|x_k^{(i)}) p(x_k^{(i)}|x_{k-1}^{(i)})}{q(x_k^{(i)}|x_{0:k-1}^{(i)}, z_{1:k})}
\]

which completes the \textit{Sequential Importance Sampling} algorithm.

\textbf{Proof:}

Indeed, the Markov property mentioned above could be written as

\[
q(x_{0:k}|z_{1:k}) = q(x_{k}|x_{0:k-1}, z_{1:k}) q(x_{0:k-1}|z_{1:k-1})
\]

We also assume that the state \( (x_k) \) is a Markov process, meaning

\[
p(x_k|x_{0:k-1}) = p(x_k|x_{k-1})
\]

and the observations \( (z_k) \) are conditionally independent given the states, so that

\[
p(z_k|x_{0:k}) = p(z_k|x_k)
\]
Finally we use the fact that at time-step $k$, all previous observations are perfectly known, and 
\[ p(z_k|x_k, z_{1:k-1}) = p(z_k|x_k) \]
Therefore
\[
\begin{align*}
   w_k(x_k) &= \frac{p(z_{1:k}|x_{0:k})p(x_{0:k})}{q(x_{0:k}|z_{1:k})} \\
   &= \frac{p(z_k|x_k)p(z_{1:k-1}|x_{0:k-1})p(x_{k-1})p(x_{0:k-1})}{q(x_k|x_{0:k-1}, z_{1:k})q(x_{0:k-1}|z_{1:k-1})}
\end{align*}
\]
which demonstrates the point. (QED)

It is important to note that the above means that the state $x_k$ cannot depend on future observations, i.e. we are dealing with Filtering and not Smoothing.

**Resampling**

One major issue with this algorithm is that the variance of the weights increases randomly over time. In order to solve this problem, we need to use a Resampling algorithm which would map our unequally weighted $x_k$’s to a new set of equally weighted sample points.

Different methods have been suggested for this. See for instance Arulampalam [14], [171].

The basic idea is to compare the Cumulative Distribution Function (CDF) created from the normalized weights, with a CDF constructed from a uniformly simulated number $U[0, 1]$. We would then eliminate the indices having too small a weight and repeat those having a sufficiently large weight.

More accurately, at a given time step $k$, for $1 \leq j \leq N_{sims}$

If 
\[
\frac{1}{N_{sims}} (U[0, 1] + j - 1) \geq \sum_{l=1}^i \tilde{w}_k^{(l)}
\]
then increment and “skip” $i$, otherwise take $x_k^{(i)}$ and set its weight to $\frac{1}{N_{sims}}$.

Note that the Resampling algorithm could create a situation where the resulting sample has many repeated points. This is known as sample impoverishment and could lead to an extreme case where all points collapse to a unique particle after a few iterations. This phenomenon is more likely if the process noise is small. One possible solution to this issue is to add a Markov-Chain Monte-Carlo (MCMC) step after the Resampling. As described further below, a Metropolis-Hastings (MH) sampling algorithm would be suitable.
Needless to say, the choice of the proposal distribution is crucial. Many suggest using
\[ q(x_k|x_{0:k-1}, z_{1:k}) = p(x_k|x_{k-1}) \]
since it will give us a simple weight identity
\[ w_k^{(i)} = w_{k-1}^{(i)} p(z_k|x_k^{(i)}) \]
Based upon this type of choices, hereafter we shall simplify and write
\[ q(x_k|x_{0:k-1}, z_{1:k}) = q(x_k|x_{k-1}, z_{1:k}) \]
without any change to our arguments.
However this choice of the proposal distribution does not take into account our most recent observation \( z_k \) at all and therefore could become inefficient.
Hence the idea of using a Gaussian Approximation for the proposal, and in particular an approximation based on the Kalman Filter, in order to incorporate the observations.
We therefore will have
\[ q(x_k|x_{k-1}, z_{1:k}) = \mathcal{N}(\hat{x}_k, P_k) \]
using the same notations as in the section on the Kalman Filter. Such filters are sometimes referred to as the Extended Particle Filter (EPF) or the Unscented Particle Filter (UPF). This is fairly similar to the *Iterative Centering Algorithm* in Kushner’s NLF.

From here, in order to estimate the parameter-set \( \Psi \) we can either use a Dual/Joint Filter, or use an ML Estimator.
Note that since the particle filter does not necessarily assume Gaussian noise, the likelihood function to be maximized has a more general form than the one used in previous sections.
Given the likelihood at step \( k \)
\[ l_k = p(z_k|z_{1:k-1}) = \int p(z_k|x_k)p(x_k|z_{1:k-1})dx_k \]
the total likelihood is the product of the \( l_k \)'s above and therefore the log-likelihood to be maximized is
\[ \ln(L_{1:N}) = \sum_{k=1}^{N} \ln(l_k) \]
(2.24)
Now \( l_k \) could be written as
\[ l_k = \int p(z_k|x_k) \frac{p(x_k|z_{1:k-1})}{q(x_k|x_{k-1}, z_{1:k})} q(x_k|x_{k-1}, z_{1:k})dx_k \]
and given that by construction the $x_k^{(i)}$'s are distributed according to $q()$, we can write the Monte-Carlo approximation

$$l_k \approx \frac{N_{\text{sims}}}{\sum_{i=1}^{N_{\text{sims}}} p(z_k|x_k^{(i)}) p(x_k^{(i)}|x_{k-1}^{(i)})}{q(x_k^{(i)}|x_{k-1}^{(i)}, z_{1:k})}$$

which we already computed for the sequential importance sampling weight update.

As we shall see in the next paragraph, it is also possible to interpret the step $k$ likelihood, as a quantity related to the total weight

$$\sum_{i=1}^{N_{\text{sims}}} w_k^{(i)}$$

Finally, we could interpret the Particle Filter as follows. We are using a Monte-Carlo simulation (via an importance sampling technique) to calculate the integral $\int f(x_k)p(x_k|z_{1:k})dx_k$.

This is exactly what other Filtering techniques try to do. The Kushner Non-Linear Filter (NLF) tries to calculate the integral via a Gaussian Quadrature. Indeed NLF uses Hermite polynomials because it treats the distributions as Normal.$^{23}$

$^{23}$Other filters cited for instance in [79] use the more general Legendre polynomials.
Implementation

Given the above theory, the algorithm for an Extended or Unscented Particle Filter could be implemented in the following way:

(1) For time step \( k = 0 \) choose \( x_0 \) and \( P_0 > 0 \). For \( i \) such that \( 1 \leq i \leq N_{sims} \) take
\[
x_0^{(i)} = x_0 + \sqrt{P_0}Z^{(i)}
\]
where \( Z^{(i)} \) is a standard Gaussian simulated number. Also take \( P_0^{(i)} = P_0 \) and
\[
w_0^{(i)} = \frac{1}{N_{sims}}
\]

While \( 1 \leq k \leq N \)

(2) For each simulation-index \( i \)
\[
\tilde{x}_k^{(i)} = KF(x_{k-1}^{(i)}) \]
with \( P_k^{(i)} \) the associated a posteriori error covariance matrix. (KF could be either the EKF or the UKF)

(3) For each \( i \) between 1 and \( N_{sims} \)
\[
\tilde{x}_k^{(i)} = \tilde{x}_k^{(i)} + \sqrt{P_k^{(i)}}Z^{(i)}
\]
where again \( Z^{(i)} \) is a standard Gaussian simulated number.

(4) Calculate the associated weights for each \( i \)
\[
w_k^{(i)} = w_{k-1}^{(i)} \frac{p(z_k|\tilde{x}_k^{(i)})p(\tilde{x}_k^{(i)}|x_{k-1}^{(i)})}{q(\tilde{x}_k^{(i)}|x_{k-1}^{(i)}, z_{1:k})}
\]
with \( q() \) the normal density with mean \( \tilde{x}_k^{(i)} \) and variance \( P_k^{(i)} \).

(5) Normalize the weights
\[
\tilde{w}_k^{(i)} = \frac{w_k^{(i)}}{\sum_{i=1}^{N_{sims}} w_k^{(i)}}
\]

(6) Resample the points \( \tilde{x}_k^{(i)} \) and get \( x_k^{(i)} \) and reset \( w_k^{(i)} = \tilde{w}_k^{(i)} = \frac{1}{N_{sims}} \).

(7) Increment \( k \), Go back to step (2) and Stop at the end of the While loop.
From step (4) we have
\[ \bar{l}_k = \sum_{i=1}^{N_{\text{sim}}} p(z_k|x_k^{(i)}) p(\hat{x}_k^{(i)}|x_{k-1}^{(i)}) q(\hat{x}_k^{(i)}|x_{k-1}^{(i)}, z_{1:k}) \]
where \( \bar{l}_k \) is a Monte-Carlo proxy for the likelihood \( l_k \) at the step \( k \). As we saw in the previous section, by minimizing
\[ -\sum_{k=1}^{N} \ln(\bar{l}_k) \]
using for instance the Direction Set algorithm, we will be maximizing the likelihood function and hence we will be obtaining the optimal parameter-set \( \hat{\Psi} \).

Given the resetting of \( w_k^{(i)} \) to a constant \( \frac{1}{N_{\text{sim}}} \) during the resampling step, we can also replace \( \bar{l}_k \) with
\[ \tilde{l}_k = \sum_{i=1}^{N_{\text{sim}}} w_k^{(i)} \]
which will provide us with an interpretation of the likelihood as the total weight.

**An Illustration**

Let us consider once again the case of the previous illustration
\[ \xi_k = \xi_{k-1} + \pi + 0.10w_k \]
and
\[ z_k = \xi_k + 0.10u_k \]
where \( \pi \approx 3.14159 \) and \( w_k, u_k \) are independent Gaussian random variables. We apply the same Kalman Filter and then apply the previous algorithm to the system. Calling
\[ n(x, m, s) = \frac{1}{\sqrt{2\pi s}} \exp \left( -\frac{(x - m)^2}{2s^2} \right) \]
the normal density with mean \( m \) and standard deviation \( s \), we will have
\[ q(\hat{x}_k^{(i)}|x_{k-1}^{(i)}, z_{1:k}) = n(\hat{x}_k^{(i)}, m = \hat{x}_k^{(i)}, s = \sqrt{P_k^{(i)}}) \]
as well as
\[ p(z_k|\hat{x}_k^{(i)}) = n(z_k, m = \hat{x}_k^{(i)}, s = 0.10) \]
and
\[ p(x_k^{(i)}|x_{k-1}^{(i)}) = n(x_k^{(i)}, m = x_{k-1}^{(i)} + \pi, s = 0.10) \]
Taking 100 particles and 500 observation points, the EPF converges very quickly to $\hat{\pi} = 3.148200$.
Alternatively the simple PF (with no Kalman component) would converge to $\hat{\pi} = 3.140266$.

Note that this example is Gaussian and Linear and therefore the Particle Filtering is \textit{not} an improvement over the Kalman Filter! Indeed the Kalman Filter is optimal for Gaussian Linear cases.

\textbf{Application to the Heston Model}

We could now apply the above Particle Filtering algorithm to our one-dimensional state where $x_k = v_k$ and $z_k = \ln S_{k+1}$ as before.

Calling
\[
n(x, m, s) = \frac{1}{\sqrt{2\pi s}} \exp \left( -\frac{(x - m)^2}{2s^2} \right)
\]
the normal density with mean $m$ and standard deviation $s$, we will have
\[
q(\tilde{x}_k^{(i)} | x_{k-1}^{(i)}, z_{1:k}) = n \left( \tilde{x}_k^{(i)}, m = \hat{x}_k^{(i)}, s = \sqrt{P_k^{(i)}} \right)
\]
as well as
\[
p(z_k | \tilde{x}_k^{(i)}) = n \left( z_k, m = z_{k-1} + (\mu_S - \frac{1}{2} \tilde{x}_k^{(i)}) \Delta t, s = \sqrt{\tilde{x}_k^{(i)} \sqrt{\Delta t}} \right)
\]
and
\[
p(\tilde{x}_k^{(i)} | x_{k-1}^{(i)}) = n \left( \tilde{x}_k^{(i)}, m = x_{k-1}^{(i)} + [\omega - \rho \xi \mu_S] - (\theta - \frac{1}{2} \rho \xi) x_{k-1}^{(i)} \frac{\Delta t}{\sqrt{\Delta t}}, s = \xi \sqrt{1 - \rho^2} \sqrt{\tilde{x}_{k-1}^{(i)} \sqrt{\Delta t}} \right)
\]
which provides us with the densities we need for the filter implementation.

The estimation of the observable state $z_k$ is
\[
\hat{z}_k = \frac{1}{N_{\text{sims}}} \sum_{i=1}^{N_{\text{sims}}} \hat{z}_k^{(i)}
\]
with $\hat{z}_k^{(i)}$ the estimation of $z_k$ from $\text{KF}(x_{k-1}^{(i)})$. 
Test Results
The results from an Extended Particle Filter (EPF) are shown in figure 2.21. The filter was constructed with the one-dimensional Heston model and was applied to a simulated time-series of 5000 points with

\[ \Psi^* = (0.40, 10.0, 0.01, -0.50) \]

As we can see in the figure, no clear superiority of the EPF is detected. The optimal parameters found via EPF are

\[ \hat{\Psi}_{EPF} = (0.020331, 0.499987, 0.040000, 0.050026) \]

which could not be considered as an improvement over

\[ \hat{\Psi}_{EKF} = (0.065886, 1.711686, 0.180884, 0.147660) \]

Again the long-term-variances are close to 0.04 for all cases, which is consistent with what we had observed.

The next natural step would be to implement and test the Unscented Particle Filter (UPF) where everything is done similarly to the EPF except the choice of the proposal distribution.

UPF has been strongly recommended by Wan and Van der Merwe in [231] and [133]. Indeed the authors claim that the filtering error from the UPF is considerably smaller than EKF, UKF or EPF.

As we can see in figure 2.22, it is true that the filtering error resulting from UPF is considerably lower than those generated from the other filters. However, the optimal parameter-set

\[ \hat{\Psi}_{UPF} = (0.020132, 0.500031, 0.040000, 0.050004) \]

obtained via UPF, is again very different from the original parameter-set \( \Psi^* \) used in the data generation.

We shall analyze the reasons behind this poor inference results more closely in the following sections.
Figure 2.21: Filtering Errors: Extended Kalman Filter and Extended Particle Filter are applied to the one-dimensional Heston model. The PF has better performance.
Figure 2.22: Filtering Errors: All Filters are applied to the one-dimensional Heston model. The PF’s have better performance.
Error Size

One possibility is that our time-series has too small an error for the filters to make a significant difference. Hence the idea of studying another case where $\Delta t = 1$ year. Let us take 200 points generated with the parameter-set

$$\Psi^* = (0.02, 0.5, 0.05, -0.5)$$

We obtain

$$\hat{\Psi}_{EKF} = (0.036, 0.093, 0.036, -1.00)$$

and

$$\hat{\Psi}_{UKF} = (0.033, 0.086, 0.033, -0.98)$$

which shows that UKF results are very close to EKF ones. Using the particle filters, we get

$$\hat{\Psi}_{EPF} = (0.019, 0.5, 0.03, -0.58)$$

which is considerably closer to the original set $\Psi^*$. Therefore EPF did bring an improvement over the traditional non-linear filters and seems to be simpler and more robust\footnote{This is because for a larger time-step the non-linearity and non-Gaussianity have a stronger impact.} than its competitors.

As for the filtering errors, it can be seen in figure 2.23 that the EPF errors are smaller than (although comparable to) those produced by EKF and UKF, which is consistent with the particle filtering theory.

As for UPF, we obtain

$$\hat{\Psi}_{UPF} = (0.019480, 0.489375, 0.047030, -0.229242)$$

which is very close to the EPF result. As we can see, the UPF errors are even smaller than those generated by EPF.

In addition to the above it would be interesting to test a Gauss-Hermite Filter (GHF) \footnote{What is more, the Girsanov theorem would not be valid and $(\xi, \rho)$ would have no reason to be the same under the risk-neutral and real measures.}. We obtain

$$\hat{\Psi}_{GHF} = (0.020398, 0.524215, 0.069661, -1.000000)$$

which is closer to the real parameter-set $\Psi^*$ compared to EKF or UKF results. However the filtering-error is more variable than those of its competitors as could be seen in figure 2.24.

Note however that this would mean that we would have access to 200 years of historic data, which is clearly unrealistic.\footnote{This issue will be revisited in the}
following sections. Also, here we generated the data via a discrete equation with $\Delta t = 1$. There was therefore no discretization error from a continuous equation. We cannot apply the same method to data coming from a continuous process.
Figure 2.23: Filters are applied to the one-dimensional Heston model. The time-series has a larger time-step $\Delta t = 1.0$. Naturally the errors are larger than the case were $\Delta t = 1/252$. 
Figure 2.24: EKF and GHF are applied to the one-dimensional Heston model. The time-series has a larger time-step $\Delta t = 1.0$. Naturally the errors are larger than the case were $\Delta t = 1/252$. 
As a measure of performance, we can compute the Mean-Price-Error (MPE) as well as the Root-Mean-Square-Error (RMSE) for each filter. These correspond respectively to the mean and the standard deviation of the plotted errors. For the MPE’s we obtain:

<table>
<thead>
<tr>
<th></th>
<th>MPE</th>
<th>RMSE</th>
</tr>
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<tbody>
<tr>
<td>EKF</td>
<td>0.007484269</td>
<td>0.003422215</td>
</tr>
<tr>
<td>UKF</td>
<td>0.007660269</td>
<td>0.003733748</td>
</tr>
<tr>
<td>GKF</td>
<td>0.009129157</td>
<td>0.005816919</td>
</tr>
<tr>
<td>EPF</td>
<td>0.007620208</td>
<td>0.002269224</td>
</tr>
<tr>
<td>UPF</td>
<td>0.007076066</td>
<td>0.001359393</td>
</tr>
</tbody>
</table>

This shows us again that the Particle Filters outperform the other ones. Again, let us remind that given 200 points with $\Delta t = 1$ and a true parameter-set $\Psi^* = (0.02, 0.5, 0.05, -0.5)$ we obtained:

<table>
<thead>
<tr>
<th></th>
<th>$\hat{\omega}$</th>
<th>$\hat{\theta}$</th>
<th>$\hat{\xi}$</th>
<th>$\hat{\rho}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>EKF</td>
<td>0.036</td>
<td>0.093</td>
<td>0.036</td>
<td>-1.00</td>
</tr>
<tr>
<td>UKF</td>
<td>0.033</td>
<td>0.086</td>
<td>0.033</td>
<td>-0.98</td>
</tr>
<tr>
<td>GKF</td>
<td>0.020</td>
<td>0.524</td>
<td>0.070</td>
<td>-1.00</td>
</tr>
<tr>
<td>EPF</td>
<td>0.019</td>
<td>0.500</td>
<td>0.033</td>
<td>-0.58</td>
</tr>
<tr>
<td>UPF</td>
<td>0.019</td>
<td>0.489</td>
<td>0.047</td>
<td>-0.22</td>
</tr>
</tbody>
</table>
The Volatility Process

The MH Enhancement

As mentioned earlier, the Resampling algorithm helps with the issue of Degeneracy, which means that it will reduce the variance of the weights. However, it might introduce a Sample Impoverishment phenomenon where all particles will have a tendency to collapse to one.

The Metropolis-Hastings (MH) algorithm could be a solution to this problem and is implemented as follows:

After the Resampling step (6) we obtain a set \( \tilde{x}_i^{(i)} \)

**(6-a)** Reapply the Kalman Filter (Extended or Unscented) to this set in order to obtain

\[ x_i^{*} = KF \left( \tilde{x}_i^{(i)} \right) \]

**(6-b)** Choose between \( x_i^{*} \) and \( \tilde{x}_i^{(i)} \) as follows:

Define

\[ \alpha = \min \left( 1, \frac{p(z_k | x_i^{*}) p(x_i^{*} | x_{i,k-1}) q(x_i^{*} | x_{i,k-1}, z_{1:k})}{p(z_k | \tilde{x}_i^{(i)}) p(\tilde{x}_i^{(i)} | x_{i,k-1}) q(\tilde{x}_i^{(i)} | x_{i,k-1}, z_{1:k})} \right) \]

then sample \( v \) from \( U[0,1] \) and

Choose \( x_i^{*} \) if \( \alpha > v \) and choose \( \tilde{x}_i^{(i)} \) if \( \alpha \leq v \).

The result is then \( x_i^{(i)} \) and we go to Step (7) as before.

Note that \( \alpha \) could be interpreted as the ratio of the non-normalized weights for the two particles we are choosing from.

Indeed

\[ \alpha = \alpha_i^{(i)} = \min \left( 1, \frac{w(x_i^{*})}{w(\tilde{x}_i^{(i)})} \right) \]

Applied to the same time-series as in the previous paragraph, the EPF with the MH modification will provide

\[ \hat{\Psi}_{EPF-MH} = (0.019, 0.499, 0.040, -0.358) \]

and

\[ MPE_{EPF-MH} = 0.007753 \]

\[ RMSE_{EPF-MH} = 0.001927 \]

compared to the previous EPF

\[ MPE_{EPF} = 0.00762 \]

\[ RMSE_{EPF} = 0.002269 \]

As we can see from the above results and the figure 2.25, there is only a marginal improvement from the introduction of the MH step in the filtering process. This is in line with the findings in the literature such as [231].
Figure 2.25: The EPF without and with the Metropolis-Hastings step is applied to the one-dimensional Heston model. The time-series has a time-step $\Delta t = 1.0$. The improvement is hardly visible.
2.3.10 Comparing Heston with other Models

We can now apply our inference tools to real market-data, in order to see which model matches the true dynamics of the assets more closely, and therefore perform Model Identification.

The Models

It is easy to generalize the Heston state space model to other stochastic volatility approaches. Indeed we could replace the Heston state equation with

\[ v_k = v_{k-1} + (\omega - \theta v_{k-1}) \Delta t + \xi v_{k-1}^{\frac{p}{2}} \sqrt{\Delta t} Z_{k-1} \]  

(2.26)

where \( p = 1/2 \) would naturally correspond to the Heston (Square-Root) model, \( p = 1 \) to the GARCH diffusion-limit model, and \( p = 3/2 \) to the \( 3/2 \) model.

These models have all been described and analyzed in [177].

The new state transition equation would therefore become

\[ v_k = v_{k-1} + \left[ \omega - \rho \xi \mu_S v_{k-1}^{\frac{p}{2}} - \left( \theta - \frac{1}{2} \rho \xi v_{k-1}^{\frac{p}{2}} \right) v_{k-1} \right] \Delta t + \rho \xi v_{k-1}^{\frac{p}{2}} \sqrt{\Delta t} \ln \left( \frac{S_k}{S_{k-1}} \right) + \xi \sqrt{1 - \rho^2} v_{k-1}^{\frac{p}{2}} \sqrt{\Delta t} Z_{k-1} \]

(2.27)

where the same choice of state space \( x_k = v_k \) is made.

For the EKF we will have

\[ A_k = 1 - \left[ \rho \xi \mu_S (p - \frac{1}{2}) v_{k-1}^{\frac{p}{2}} + \theta - \frac{1}{2} \rho \xi (p + \frac{1}{2}) v_{k-1}^{\frac{p}{2}} \right] \Delta t + (p - \frac{1}{2}) \rho \xi v_{k-1}^{\frac{p}{2}} \ln \left( \frac{S_k}{S_{k-1}} \right) \]

and

\[ W_k = \xi \sqrt{1 - \rho^2} v_{k-1}^{\frac{p}{2}} \sqrt{\Delta t} \]

as well as

\[ H_k = -\frac{1}{2} \Delta t \]

and

\[ U_k = \sqrt{v_k} \sqrt{\Delta t} \]

The same time update and measurement update equations could be used with the UKF or Kushner’s NLF.

We could also apply the Particle Filtering algorithm to our problem. Using the same notations as in section 1.3.2 and calling

\[ n(x, m, s) = \frac{1}{\sqrt{2\pi}s} \exp \left( -\frac{(x - m)^2}{2s^2} \right) \]

the Normal density with mean \( m \) and standard deviation \( s \), we will have

\[ q(\tilde{x}_k^{(i)}|x_{k-1}^{(i)}, z_{1:k}) = n \left( \tilde{x}_k^{(i)}, m = \tilde{x}_k^{(i)}, s = \sqrt{r_k^{(i)}} \right) \]
as well as

\[ p(z_k|x_k^{(i)}) = n \left( z_k, m = z_{k-1} + (\mu S - \frac{1}{2} \tilde{x}_k^{(i)}) \Delta t, s = \sqrt{\tilde{x}_k^{(i)} \Delta t} \right) \]

and

\[ p(\tilde{x}_k^{(i)}|x_{k-1}^{(i)}) = n \left( \tilde{x}_k^{(i)}, m_x, s = \xi \sqrt{1 - \rho^2 (x_{k-1}^{(i)})^p \Delta t} \right) \]

with

\[ m_x = x_{k-1}^{(i)} + \left[ \omega - \rho \xi \mu S (x_{k-1}^{(i)})^p - \frac{1}{2} \rho \xi (x_{k-1}^{(i)})^p \tilde{x}_k^{(i)} \Delta t + \rho \xi (x_{k-1}^{(i)})^p \Delta t \right] \]

and as before we have

\[ u_k^{(i)} = u_{k-1}^{(i)} \frac{p(z_k|\tilde{x}_k^{(i)})p(\tilde{x}_k^{(i)}|x_{k-1}^{(i)})}{q(\tilde{x}_k^{(i)}|x_{k-1}^{(i)}, z_{1:k})} \]

which provides us with what we need for the filter implementation.

**The Results**

The above filters were applied to five years of S&P500 time-series (1996 to 2001) and the filtering errors were considered for the Heston model, the GARCH model and the 3/2 model. Daily index close-prices were used for this purpose, and the time interval was set to \( \Delta t = 1/252 \).

<table>
<thead>
<tr>
<th>Filter</th>
<th>MPE</th>
<th>RMSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>EKF-Heston</td>
<td>3.58207e-05</td>
<td>1.83223e-05</td>
</tr>
<tr>
<td>EKF-GARCH</td>
<td>2.78438e-05</td>
<td>1.42428e-05</td>
</tr>
<tr>
<td>EKF-3/2</td>
<td>2.63227e-05</td>
<td>1.74760e-05</td>
</tr>
<tr>
<td>UKF-Heston</td>
<td>3.00000e-05</td>
<td>1.91280e-05</td>
</tr>
<tr>
<td>UKF-GARCH</td>
<td>2.99275e-05</td>
<td>2.58131e-05</td>
</tr>
<tr>
<td>UKF-3/2</td>
<td>2.82279e-05</td>
<td>1.55777e-05</td>
</tr>
<tr>
<td>EPF-Heston</td>
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<td>1.34534e-05</td>
</tr>
<tr>
<td>EPF-GARCH</td>
<td>2.48733e-05</td>
<td>4.99337e-06</td>
</tr>
<tr>
<td>EPF-3/2</td>
<td>2.26462e-05</td>
<td>2.58645e-06</td>
</tr>
<tr>
<td>UPF-Heston</td>
<td>2.04000e-05</td>
<td>2.74818e-06</td>
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<td>UPF-GARCH</td>
<td>2.69036e-05</td>
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</tr>
<tr>
<td>UPF-3/2</td>
<td>1.73857e-05</td>
<td>4.09918e-06</td>
</tr>
</tbody>
</table>
Figure 2.26: Comparison of EKF Filtering errors for Heston, GARCH and $3/2$ Models. The latter seems to perform better.
Figure 2.27: Comparison of UKF Filtering errors for Heston, GARCH and $3/2$ Models. The latter seems to perform better.
Figure 2.28: Comparison of EPF Filtering errors for Heston, GARCH and $3/2$ Models. The latter seems to perform better.
Figure 2.29: Comparison of UPF Filtering errors for Heston, GARCH and \( \frac{3}{2} \) Models. The latter seems to perform better.
Figure 2.30: Comparison of Filtering errors for the Heston Model. PF’s seem to perform better.
Figure 2.31: Comparison of Filtering errors for the GARCH Model. PF’s seem to perform better.
Figure 2.32: Comparison of Filtering errors for the 3/2 Model. PF's seem to perform better.
Two immediate observations can be made: On the one hand the Particle Filters have a better performance than the Gaussian ones, which reconfirms what one would anticipate. On the other hand for most of the Filters, the $3/2$ model seems to outperform the Heston model, which is in line with the findings of Engle & Ishida [95]. Again, this shows that the filtering process could be used not only for Parameter-Estimation but also for Model Identification.

This suggests further filtering on other existing models such as Jump Diffusion [190]. Clearly, because of the non-Gaussianity of jump based models, the Particle Filtering technique will need to be applied to them.

**Parameter Learning Revisited**

We tried a JF via the Kalman Filter where the parameters were given a prior distribution. We can now apply the Particle Filtering techniques to this framework as in [176], [224]:

We simulate $x_k^{(i)}$ at time-step $k$ from the prior $p(x_k | x_{k-1}^{(i)})$ and we also simulate each parameter $\psi^{(i)}$ from its prior $q(\psi) = \mathcal{N}(m_\psi, s_\psi)$ where these mean and standard-deviations are to be determined.

We then update the priors by incorporating the observation $z_k$

$$p(x_k | z_k) \propto p(z_k | x_k^{(i)}, \psi^{(i)})p(x_k | x_{k-1}^{(i)})$$

and similarly

$$p(\psi | z_k) \propto p(z_k | x_k^{(i)}, \psi^{(i)})p(\psi | x_{k-1}^{(i)})$$

and we obtain the posterior distributions.

Calling

$$w_k^{(i)} = \frac{p(z_k | x_k^{(i)}, \psi^{(i)})w_{k-1}^{(i)}}{\sum_{i=1}^{N_{sims}} p(z_k | x_k^{(i)}, \psi^{(i)})w_{k-1}^{(i)}}$$

We now have the posteriors of $x_k$ and $\psi$ and we can simulate them for the following step via a Metropolis-Hastings (MH) accept/reject technique with the proposal distribution $q(\psi, m_\psi, s_\psi)$ with

$$m_\psi = \sum_{i=1}^{N_{sims}} w_k^{(i)} \psi^{(i)}$$

and

$$s_\psi = \sum_{i=1}^{N_{sims}} w_k^{(i)} (\psi^{(i)} - m_\psi)^2$$

The MH step will consist in the following: we accept the simulation point $\tilde{\psi}^{(i)}$ from $q(\psi)$ with a probability $\alpha(\psi^{(i)}, \tilde{\psi}^{(i)})$ where $\forall i$ between 1 and $N_{sims}$ we have

$$\alpha(\psi^{(i)}, \tilde{\psi}^{(i)}) = \min \left(1, \frac{p(\tilde{\psi}^{(i)} | z_k)q(\psi^{(i)})}{p(\psi^{(i)} | z_k)q(\tilde{\psi}^{(i)})} \right)$$
In practice we simulate a uniform random-variable $u$ and accept the simulated point $\tilde{\psi}^{(i)}$ if $\alpha > u$, and reject it (and keep $\psi^{(i)}$) otherwise.

We keep simulating alternatively the state variable and each parameter by incorporating the observations at each step and wait for the parameters to converge to their ideal mean.

It is important to note that this Joint Filtering differs from the usual MCMC techniques such as [156], [92] where we update the particles by incorporating all observations at each simulation-step.
2.3.11 The Performance of the Inference Tools

We have applied various Gaussian and Particle-based Filters to daily historic data. None of the methodologies performed very well at that frequency. We now try to analyze the reasons behind this issue.

A known weakness of optimization algorithms is the following. The higher the number of parameters, the worse the performance of the algorithm. This means that a one-parameter optimization should perform best. To test this, we simulate 5000 points via the Heston model with a parameter-set $\Psi^*$ as shown below (Also see figure 2.33).

We use a drift of $\mu_S = 0.025$ and a time-step $\Delta t = 1/252$ as before. In order to get the best performance we fix all parameters except one. For instance to obtain $\hat{\omega}$ we fix $\theta = 10.0, \xi = 0.03, \rho = -0.50, \mu_S = 0.025$, we choose a reasonable initial point $\omega_0$ and then optimize upon $\omega$ only. We choose an initial parameter-set $\Psi_0$ as shown below. The results are displayed in the table 2.4.

It is interesting to note that the estimation of the volatility-drift parameters $\Psi^*$

<table>
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<tr>
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<th>$\hat{\xi}$</th>
<th>$\hat{\rho}$</th>
</tr>
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<tr>
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</tr>
<tr>
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<td>0.000001</td>
<td>+0.598434</td>
</tr>
<tr>
<td>EPF</td>
<td>0.098287</td>
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<td>0.044437</td>
<td>-0.827729</td>
</tr>
<tr>
<td>UPF</td>
<td>0.100581</td>
<td>10.221816</td>
<td>0.051902</td>
<td>-0.487695</td>
</tr>
</tbody>
</table>

Table 2.4: The optimal parameter-set $\hat{\Psi}$. The estimation is performed individually for each parameter on the artificially generated time-series. Particle Filters use 1000 simulations.

Note that in this section we are not checking the validity of the assumption that the real stock-market follows a Heston (or another) process. We assume we know the process exactly, and try to recover the embedded parameters.

We made the 5000 daily simulations directly from the discretized SDE with a $\Delta t = 1/252$. We also tried simulating 5,000,000 points with $\Delta t = 1/252,000$ and sampling 5000 daily points from there. Although the second method is more correct, the difference in results was small, which means that the Euler discretization is sufficiently accurate at the daily level. This is in agreement with results found by Elerian [92].
The estimation of volatility and correlation parameters \((\xi, \rho)\) is not as straightforward. This could be seen by plotting the likelihood \(L(\Psi)\) as a function of \(\omega, \theta, \xi\) and \(\rho\) separately. We fix three parameters to their optimal values and plot \(L(\Psi)\) as a function of the last one. We observe in figures 2.34 to 2.37 that the likelihood function is fairly easy to optimize for \((\omega, \theta)\). However the function is very flat around the optimal \(\xi\) and \(\rho\). Hence the difficulty of finding the optimums!

\[^{28}\text{A joint estimation of } (\omega, \theta) \text{ based on the same data-set with known } (\xi, \rho) \text{ provides } (\hat{\omega} = 0.117889, \hat{\theta} = 11.996760)\]

(\(\omega, \theta\)) could be done fairly well via EKF.\(^{28}\) This makes sense since the dependence on these parameters is linear.
Figure 2.33: Simulated Stock-Price path via Heston using \( \Psi^* \). This is an artificial time-series following the Heston model.
Figure 2.34: $f(\omega) = L(\omega, \hat{\theta}, \hat{\xi}, \hat{\rho})$ has a good slope around $\hat{\omega}=0.10$. 
Figure 2.35: $f(\theta) = L(\hat{\omega}, \hat{\theta}, \hat{\xi}, \hat{\rho})$ has a good slope around $\hat{\theta}=10.0$. 
Figure 2.36: \( f(\xi) = L(\hat{\omega}, \hat{\theta}, \xi, \hat{\rho}) \) is flat around \( \hat{\xi} = 0.03 \).
Figure 2.37: $f(\rho) = L(\hat{\omega}, \hat{\theta}, \hat{\xi}, \rho)$ is flat and irregular around $\rho=-0.50$. 
Sample Size

It seems therefore that the estimation is inefficient for the parameter $\xi$ no matter which filter we use. The issue is that of inefficiency (large error variance) for this given sample size. This is indeed one of the shortcomings of the Maximum Likelihood Estimators (MLE). For a given sample size they can very well be inefficient and even have a bias. The choice of the filter will not solve this issue.

However (under minimal regularity conditions) MLE’s are consistent and therefore asymptotically converge to the correct optimum. This means that the sample size is key. To test this we can choose larger samples of $N = 50,000$, $N = 100,000$ and $N = 500,000$ points and rerun the simplest filter, namely the EKF. As expected the optimum of the Likelihood function becomes closer and closer to $\xi^\star$. This can be seen in figures 2.38 to 2.41 as well as in table 2.5.

Same exact observations could be made for the correlation parameter $\rho$ and the results are displayed in the same table 2.5. The likelihood graphs are omitted in the interest of brevity.

As for the drift parameters $\omega$ and $\theta$, the convergence was good even for $N = 5000$ as previously observed.

Unfortunately in reality we have limited historic data. Even at a daily frequency 50,000 points would correspond to 200 years!

One possibility would be to use intra-day data, however that assumes that the behavior of the stock-price is the same intra-day (which is reasonable considering we started with a continuous SDE). Moreover, clean intra-day data is usually not readily available and needs preprocessing.

<table>
<thead>
<tr>
<th>$N$</th>
<th>$\hat{\xi}$</th>
<th>$\hat{\rho}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>5000</td>
<td>0.032324</td>
<td>-0.873571</td>
</tr>
<tr>
<td>50,000</td>
<td>0.034663</td>
<td>-0.608088</td>
</tr>
<tr>
<td>100,000</td>
<td>0.033400</td>
<td>-0.556868</td>
</tr>
<tr>
<td>500,000</td>
<td>0.031922</td>
<td>-0.532142</td>
</tr>
</tbody>
</table>

Table 2.5: The Optimal EKF parameters $\hat{\xi}$ and $\hat{\rho}$ given a sample size $N$.

The true parameters are $\xi^\star = 0.03$ and $\rho^\star = -0.50$.

The initial values were $\xi_0 = 0.02$ and $\rho_0 = -0.40$.

Therefore, having $p$ parameters in the optimal parameter-set $\hat{\Psi}_N = \left( \hat{\Psi}_N[j] \right)_{1 \leq j \leq p}$

---

29 A known and simple example for the bias of MLEs, is that of the estimation of the variance of a Gaussian sequence of a finite size $(x_1, ..., x_N)$. The ML estimate for the mean is $\hat{\mu}_N = \frac{1}{N} \sum_{k=1}^{N} x_k$ and the ML estimate for the variance is $\hat{\sigma}_N = \frac{1}{N} \sum_{k=1}^{N} (x_k - \hat{\mu}_N)^2$. The latter ML estimation is biased and the correct estimation would be $\hat{\sigma}_N = \frac{1}{N-1} \sum_{k=1}^{N} (x_k - \hat{\mu}_N)^2$. However it is clear that as $N \to +\infty$ we have $\hat{\sigma}_N \approx \hat{\sigma}_N$ and the bias gradually disappears.
for a sample size $N$, we have for each parameter $\Psi[j]$

$$\lim_{N \to +\infty} \hat{\Psi}_N[j] | \{\Psi[k] = \Psi^*[k]; 1 \leq k \leq p; k \neq j\} = \Psi^*[j] \quad (2.28)$$

What is more, this is true for any valid initial value $\Psi_0[j]$, which means the MLE is robust.
Figure 2.38: $f(\xi) = L(\hat{\omega}, \hat{\theta}, \hat{\rho})$ via EKF for $N = 5000$ points. The true value is $\xi^* = 0.03$. 
Figure 2.39: $f(\xi) = L(\hat{\omega}, \hat{\theta}, \hat{\rho})$ via EKF for $N = 50,000$ points. The true value is $\xi^* = 0.03$. 
Figure 2.40: $f(\xi) = L(\omega, \theta, \xi, \rho)$ via EKF for $N = 100,000$ points. The true value is $\xi^* = 0.03$. 
Figure 2.41: $f(\xi) = L(\hat{\omega}, \hat{\theta}, \xi, \hat{\rho})$ via EKF for $N = 500,000$ points. The true value is $\xi^* = 0.03$. 
Joint Estimation of the Parameters

Let us now assume that we do not know any of the parameters, choose an initial set \( \Psi_0 \) and test the consistency of the MLE. We shall apply the EKF to the data and take the same true parameter set \( \Psi^* \) as in the previous section. We assume that \( \mu_S = 0.025 \) is known, otherwise it could be estimated together with the model parameters.

As previously mentioned, the likelihood function becomes flat and therefore harder to maximize under a higher number of parameters. The convergence of the estimator will therefore be slower.

Despite this, we can observe in table 2.8 the asymptotic convergence of the estimator even under the joint estimation of all parameters. Indeed we have now

\[
\lim_{N \to +\infty} \hat{\Psi}_N = \Psi^*
\]  

(2.29)

which corresponds to the generalization of (2.28) in the previous section.

\[
\begin{array}{c|c|c|c|c|c|c}
\Psi^* & \omega^* & \theta^* & \xi^* & \rho^* \\
0.10 & 10.0 & 0.03 & -0.50 \\
\end{array}
\]

Table 2.6: The true parameter-set \( \Psi^* \) used for data generation.

\[
\begin{array}{c|c|c|c|c|c|c}
\Psi_0 & \omega_0 & \theta_0 & \xi_0 & \rho_0 \\
0.15 & 15.0 & 0.02 & -0.40 \\
\end{array}
\]

Table 2.7: The initial parameter-set \( \Psi_0 \) used for the optimization process.

\[
\begin{array}{c|c|c|c|c|c|c}
N & \hat{\omega} & \hat{\theta} & \hat{\xi} & \hat{\rho} \\
5000 & 0.150854 & 15.294576 & 0.266175 & -0.128835 \\
50,000 & 0.126387 & 12.748852 & 0.020521 & -1.000000 \\
100,000 & 0.136023 & 13.700906 & 0.044353 & -0.439961 \\
500,000 & 0.100097 & 10.030336 & 0.061688 & -0.257305 \\
1,000,000 & 0.105264 & 10.548642 & 0.043818 & -0.356234 \\
2,000,000 & 0.103183 & 10.334876 & 0.039767 & -0.374677 \\
4,000,000 & 0.105292 & 10.538019 & 0.043288 & -0.347562 \\
5,000,000 & 0.101097 & 10.118951 & 0.028588 & -0.514346 \\
\end{array}
\]

Table 2.8: The Optimal EKF parameter-set \( \hat{\Psi} \) for given a sample size \( N \). The four parameters are estimated jointly.
We ran other Filters (UKF, EPF, UPF) on the same data set and observed only marginal improvement. The results are omitted in the interest of brevity. It therefore seems that the fundamental issue is related to the slow convergence of the MLEs regardless of the Filtering method.

A related issue previously mentioned is the size of the observation error $U_k \propto \sqrt{\Delta t}$ which is large compared to the observation function $H_k \propto \Delta t$ for daily observations.
**Error Size revisited**

As previously mentioned, this underlines the more fundamental problem for the SV Estimation: By definition, volatility represents the noise of the stock process. Indeed if we had taken the spot price $S_k$ as the observation and the variance $v_k$ as the state, we would have

$$S_{k+1} = S_k + S_k \mu \Delta t + S_k v_k \sqrt{\Delta t} B_k$$

we would then have an observation function gradient $H = 0$ and the system would be **unobservable**!

It is precisely because we use a Taylor second order expansion

$$\ln(1 + x) \approx x - \frac{1}{2} x^2$$

that we obtain access to $v_k$ through the observation function. However in

$$\ln \left( \frac{S_{k+1}}{S_k} \right) = (\mu - \frac{1}{2} v_k) \Delta t + \sqrt{v_k} \sqrt{\Delta t} B_k$$

the error remains **dominant** as the first order of the expansion.\(^{30}\)

Harvey, Ruiz and Shephard [130] use the approximation $\Delta t = o(\sqrt{\Delta t})$ and take

$$z_k = \ln \left( \ln^2 \left( \frac{S_{k+1}}{S_k} \right) \right) \approx \ln(v_k) + \ln(\Delta t) + \ln(B_k^2)$$

Note that under this form EKF would blow up since $z_k^- = h(v_k, 0) = -\infty$.

They therefore use the fact that $\mathbb{E}[\ln(B_k^2)] = -1.27$ and stdev$[\ln(B_k^2)] = \pi/\sqrt{2}$

and consider the Gaussian approximation

$$\ln(B_k^2) \sim -1.27 + \frac{\pi}{\sqrt{2}} N(0, 1)$$

which may or may not be valid. We call this approximation Harvey-Ruiz-Shephard (HRS) and apply it to the same case as in the previous paragraphs.

As can be seen in table 2.9 the approximation seems to be valid for our example. Note that UKF would not have this issue since we would work with the real nonlinear function $z = h(x, u)$ above. However we would still deal with log’s of very small quantities which could be numerically unstable.

\(^{30}\) Note that this is different from a Variance Swap where we work with the **expected values**. Indeed the approximation is perfectly valid if for the return $R = \Delta S/S$ we write

$$\mathbb{E}[\ln(1 + R) - R] \approx -\frac{1}{2} v$$

but again, the approximation breaks if we work for one sample path.
Another way of tackling the same equation would be via a Particle Filter where

$$z_k = \ln \left( \left| \ln \left( \frac{S_{k+1}}{S_k} \right) \right| \right) \approx \frac{1}{2} \ln(v_k) + \frac{1}{2} \ln(\Delta t) + \ln(|B_k|)$$

and as stated in [10] the density of $\ln(|B_k|)$ is

$$f(x) = 2e^x n(e^x)$$

with $n()$ the normal density.\(^{31}\)

Testing the same data set provides table 2.10 which does not seem to improve upon the KF.

<table>
<thead>
<tr>
<th>$N$</th>
<th>$\hat{\omega}$</th>
<th>$\hat{\theta}$</th>
<th>$\hat{\xi}$</th>
<th>$\hat{\rho}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>5000</td>
<td>0.722746</td>
<td>71.753861</td>
<td>0.044602</td>
<td>-1.000000</td>
</tr>
<tr>
<td>50,000</td>
<td>0.234110</td>
<td>23.575193</td>
<td>0.028056</td>
<td>-1.000000</td>
</tr>
<tr>
<td>100,000</td>
<td>0.150512</td>
<td>15.186113</td>
<td>0.017748</td>
<td>-1.000000</td>
</tr>
<tr>
<td>500,000</td>
<td>0.109738</td>
<td>11.020391</td>
<td>0.027140</td>
<td>-0.531481</td>
</tr>
</tbody>
</table>

Table 2.9: The Optimal EKF parameter-set $\hat{\Psi}$ via the HRS approximation for given a sample size $N$. The four parameters are estimated jointly.

Table 2.10: The Optimal PF parameter-set $\hat{\Psi}$ for given a sample size $N$. The four parameters are estimated jointly.

\(^{31}\)It is easy to see that if $X$ is a standard Normal variable, then the CDF of $\ln(|X|)$ is

$$F(x) = P(\ln(|X|) \leq x) = P(|X| \leq e^x) = P(-e^x \leq X \leq e^x)$$

therefore

$$F(x) = N(e^x) - N(-e^x) = 2N(e^x) - 1$$

and the density is determined by taking the derivative with respect to $x$ as usual.
It is important to note that even if we took the example of the Heston model, the same issues are true for any stochastic volatility model of type

$$v_k = v_{k-1} + (\omega - \theta v_{k-1})\Delta t + \xi v_{k-1}^p \sqrt{\Delta t} Z_{k-1}$$

even including the GARCH-diffusion and the 3/2 models.

Indeed, as previously mentioned, even if the Transition equation is different here, the Observation equation remains the same.

Applying the EKF, we have the Transition matrix and noise

$$A_k = 1 - \left[ \rho \xi \mu_s (p - \frac{1}{2}) v_{k-1}^{p-1} + \theta - \frac{1}{2} \rho \xi (p + \frac{1}{2}) v_{k-1}^{p-1} \right] \Delta t + (p-\frac{1}{2}) \rho \xi v_{k-1}^{p-1} \ln \left( \frac{S_k}{S_{k-1}} \right)$$

$$W_k = \xi \sqrt{1 - \rho^2 v_{k-1}^p \sqrt{\Delta t}}$$

However, we still have the Observation matrix and noise

$$H_k = -\frac{1}{2} \Delta t$$

and

$$U_k = \sqrt{v_k} \sqrt{\Delta t}$$

and the same issue of $\Delta t = o(\sqrt{\Delta t})$ still exists at observation level for any value of $p$.

Another point that should be mentioned is that even if $\xi$ and $\rho$ are separately harder to estimate than $\omega$ and $\theta$, the product $\rho \xi$ appears in the equations at the same level. Indeed as we just saw, in $A_k$ only the product $\rho \xi$ is available. However at the noise level $W_k$, we can distinguish the two parameters $\rho$ and $\xi$.

For instance, in our previous EKF joint estimation table we had for 50,000 points $\xi \approx 0.020521$, $\rho \approx -1.0000$ and again, the individual estimations of $\xi$ and $\rho$ remained far from their true values. However we have $\hat{\xi} \hat{\rho} \approx -0.020521$ which is much closer to $\xi^* \rho^* = -0.015$.

Interestingly, the product $\rho \xi$ is what we need to determine the Skewness of the distribution.\(^{32}\) However, we do need to determine $\xi$ alone to obtain the distribution Kurtosis.

It is also worth noting that in a GARCH framework, we do not have this issue of poor Observability for the discrete case. Indeed at each point in time, $v_k$ is known exactly as a function of previous observations. Only later, we go to the two-factor Diffusion limit as Nelson [194] does.

However, we have to bear in mind that this GARCH Diffusion limit is a very special case of the Stochastic Volatility problem, since it misses the second source of randomness in the discrete case. As Corradi [61] explains, a discrete GARCH model may very well converge towards a one-factor diffusion process without stochastic volatility. Interestingly, when discretizing the one-factor continuous

\(^{32}\)This remark will be used in the following chapter.
process we can recover GARCH, whereas when discretizing the two-factor continuous process we will not obtain GARCH but the two-factor discrete process we have been working with. This explains why GARCH MLE (without Filtering) can recover parameters used in a simulated time-series of length 5000 created via a one-factor GARCH process, while it cannot recover the diffusion-limit parameters from a time-series created via a two-factor Stochastic Volatility process as accurately. Indeed, one could see this in tables 2.11 and 2.12.

<table>
<thead>
<tr>
<th>$\Psi^*$</th>
<th>$\omega$</th>
<th>$\alpha$</th>
<th>$\beta$</th>
<th>$c$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.00000176</td>
<td>0.0626</td>
<td>0.89760</td>
<td>0.00</td>
</tr>
<tr>
<td>$\Psi$</td>
<td>0.00000200</td>
<td>0.0530</td>
<td>0.89437</td>
<td>0.05</td>
</tr>
</tbody>
</table>

Table 2.11: Real and Optimal parameter-sets obtained via NGARCH MLE. The 5000 points were generated via the one-factor NGARCH with daily parameters

<table>
<thead>
<tr>
<th>$\Psi^*$</th>
<th>$\omega$</th>
<th>$\theta$</th>
<th>$\xi\rho$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.100000</td>
<td>10.00</td>
<td>-0.015</td>
</tr>
<tr>
<td>$\Psi_{GARCH}$</td>
<td>0.063504</td>
<td>6.84</td>
<td>-0.019</td>
</tr>
<tr>
<td>$\Psi_{EKF}$</td>
<td>0.148000</td>
<td>14.48</td>
<td>-0.023</td>
</tr>
</tbody>
</table>

Table 2.12: Real and Optimal parameter-sets obtained via NGARCH MLE as well as EKF. The 5000 points were generated via the two-factor GARCH Diffusion limit with annual parameters

This also explains why estimating $\omega$ and $\theta$ alone works so much better with 5000 points. After all if we had $\xi = 0$ and therefore a deterministic instantaneous variance, we would have no observability problem to talk about. Indeed

\[ \Delta t = o(\sqrt{\Delta t}) \]

Indeed it would be tempting to try to get around the $\Delta t = o(\sqrt{\Delta t})$ issue by rewriting the equations via daily parameters $\mu_S^d = \mu_S \Delta t$ and $v_k^d = \Delta t v_k$ as well as $\omega^d = \Delta t^2 \omega$, $\theta^d = \Delta t \theta$ and $\xi^d = \Delta t \xi$ with $\rho$ remaining unchanged. Dropping the superscript $d$ for simplifying the notations, we shall have

\[
\ln S_{k+1} = \ln S_k + \mu_S - \frac{1}{2} v_k + \sqrt{v_k} B_k \\
v_{k+1} = v_k + \omega - \theta v_k + \xi \sqrt{v_k} Z_k
\]

which seems to have eliminated the difficulty. However now we have

\[ v_k = o(\sqrt{v_k}) \]

which was not the case with yearly variances, and the same poor observability issue arises again! We therefore see that the heart of the difficulty is a low signal-to-noise ratio (SNR) for the problem at hand.
$v_t$ would be exactly known at each time-step, as is the case in a GARCH framework.

Finally, we can now see better why the estimation worked fairly well even with 200 points if $\Delta t = 1$ year. Simply because we do not have $\Delta t = o(\sqrt{\Delta t})$ and the observability is much more accurate. Having said that, with such a large $\Delta t$ other issues such as strong non-linearity and the non-applicability of the Grisanov theorem arise. Not to mention the fact that 200 points would correspond to two-hundred years of data!
**High Frequency Data**

Given that the results seem to converge for a large number of data points, one idea would be to use a higher sampling frequency. Indeed if instead of using daily data we sample every five seconds, one a ten year range we will have $10 \times 252 \times 6.5 \times 60 \times 60 \div 5 = 11,793,600$ data points which is very sufficient for our MLE’s.

For testing the use of high frequency data, we can generate via Monte-Carlo 5,000,000 points with a $\Delta t = 1/252,000$ which corresponds to 20 years. We obtain the results in the table 2.13 below. Both rows have reasonable results. It is however notable that the EKF/ HRS method seems to perform better than the plain EKF.

<table>
<thead>
<tr>
<th></th>
<th>$\hat{\omega}$</th>
<th>$\hat{\theta}$</th>
<th>$\hat{\xi}$</th>
<th>$\hat{\rho}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>EKF</td>
<td>0.090280</td>
<td>9.019962</td>
<td>0.042984</td>
<td>-0.283236</td>
</tr>
<tr>
<td>EKF/ HRS</td>
<td>0.092372</td>
<td>9.224421</td>
<td>0.030951</td>
<td>-0.507763</td>
</tr>
</tbody>
</table>

Table 2.13: The Optimal parameter-set $\hat{\Psi}$ for 5,000,000 data points. The sampling is performed 1000 times a day and therefore the data-set corresponds to 5000 business days. The four parameters are estimated jointly.

It may seem a little surprising that for the same time period $[0, T]$ dividing $\Delta t$ by 1000 and multiplying $N$ by 1000 helps us. Indeed, why don’t the two operations cancel one another?

However, observing the negative of log-Likelihood function in an EKF framework

$$\phi(\omega, \theta, \xi, \rho) = \sum_{k=1}^{N} \left[ \ln(F_k) + \frac{z_k^2}{F_k} \right]$$

with

$$\tilde{z}_k = z_k - h(\tilde{x}_k^-)$$

and

$$F_k = H_k P_k^- H_k^t + U_k U_k^t$$

We can see that considering first order terms, dividing $\Delta t$ by 1000, or equivalently multiplying it by $\epsilon = 1/1000$, will cause the transition matrix $A_k$ to be unchanged, the transition noise $W_k$ to be multiplied by $\sqrt{\epsilon}$, the observation matrix $H_k$ to be multiplied by $\epsilon$ and the observation noise $U_k$ to be multiplied by $\sqrt{\epsilon}$.

What is more $A_k$ being unchanged, will cause $P_k^-$ and $P_k$ to remain unchanged as well. Therefore $\tilde{z}_k$ will be multiplied by $\sqrt{\epsilon}$ and the term $F_k$ will be multiplied by $\epsilon$ and the fraction used in the log-Likelihood sum will remain the same. This causes the sum $\phi(\omega, \theta, \xi, \rho)$ to be higher by a factor $1/\epsilon$ which shows that higher frequency does allow us to obtain a higher Likelihood function and therefore
better convergence. This is in agreement with what we observed in the above test.

The Frequency of the Observations

Note that the ideal stochastic Differential Equations are supposed to be continuous, however we only have discrete observations obtained via an Euler scheme. This introduces a discretization Error which may become important as the time-interval $\Delta t$ becomes larger.

As mentioned in [92], [164], [201] the solution would be to fill the missing data via additional simulations in time:

For the observation time-step $1 \leq k \leq N$, the simulation $1 \leq i \leq N_{sims}$ and the additional time-step $1 \leq j \leq M$ we would have the particles

$$
\tilde{x}_{k+\frac{j}{M}}^{(i)} = \tilde{x}_{k+\frac{j-1}{M}}^{(i)} + (\omega - \theta \tilde{x}_{k+\frac{j-1}{M}}^{(i)}) \frac{\Delta t}{M} + \xi \sqrt{\tilde{x}_{k+\frac{j-1}{M}}^{(i)}} \sqrt{\frac{\Delta t}{M}} Z_{k+\frac{j}{M}}^{(i)}
$$

and the observation

$$
z_{k+1} = z_k + \sum_{j=1}^{M} (\mu S - \frac{1}{2} \tilde{x}_{k+\frac{j}{M}}) \frac{\Delta t}{M} + \sqrt{\sum_{j=1}^{M} \tilde{x}_{k+\frac{j}{M}} \Delta t} B_k
$$

where each $Z_{k+\frac{j}{M}}^{(i)}$ has a correlation $\rho$ with $B_k$. Naturally the innovations $Z_t$ are mutually uncorrelated.

However as discussed in [164] the discretization error is small when $\Delta t = 1/252$, which is the case we are dealing with.
Sampling Distribution

Even if in practice we deal with one historic path, we should determine the distribution of the optimal parameter-set as follows. We simulate $P = 500$ paths of length $N = 5000$ and estimate for each path $j$ the optimal set $\hat{\Psi}^{(j)}$. We then can estimate

$$\tilde{\Psi} = \frac{1}{P} \sum_{j=0}^{P-1} \hat{\Psi}^{(j)}$$

as well as the variance

$$V(\tilde{\Psi}) = \frac{1}{P} \sum_{j=0}^{P-1} (\hat{\Psi}^{(j)} - \tilde{\Psi})^2$$

This way we will know how the estimator performs in average and how far we could be from this average. The distribution of the parameter-set around its mean is referred to as the Sampling Distribution [168].

As we can see in table 2.14 the average-estimated parameter-set is closer to the true-set than the one-path-estimated set we were considering in the previous section. However the corresponding standard-deviation is quite high and we could very well get poor results as previously seen.

From figures 2.42 to 2.45 we can see that for this data length $N$ and this sample size $P$ the parameters $\omega$ and $\theta$ are determined via EKF in a fairly unbiased way. However the estimator is not efficient and has a large standard-deviation. As for $\xi$ and $\rho$ we have both bias and inefficiency.

This is not surprising given the results of the previous paragraphs. We obtained good results for $(\omega, \theta)$ when estimated alone, and not so good results for $(\xi, \rho)$. As mentioned, classical filtering theory works well when the parameters affect the drift of the observation and not the noise. This causes a slow convergence issue for all our parameters. But this is doubly true for $(\xi, \rho)$ since they affect the “noise of the noise”.

As previously observed the bias and inefficiency will disappear as $N \to +\infty$ as is the case for any MLE estimator. Indeed the biases and the standard-deviations are smaller for $N = 50,000$ than for $N = 5000$ as we can see in table 2.14.
Figure 2.42: Density for $\hat{\omega}$ estimated from 500 paths of length 5000 via EKF. The true value is $\omega^* = 0.10$. The sampling distribution is fairly unbiased, but is inefficient.
Figure 2.43: Density for $\hat{\theta}$ estimated from 500 paths of length 5000 via EKF. The true value is $\theta^* = 10$. The sampling distribution is fairly unbiased, but is inefficient.
Figure 2.44: Density for $\hat{\xi}$ estimated from 500 paths of length 5000 via EKF. The true value is $\xi^* = 0.03$. The sampling distribution is inefficient and even has a bias.
Figure 2.45: Density for $\hat{\rho}$ estimated from 500 paths of length 5000 via EKF. The true value is $\rho^* = -0.50$. The sampling distribution is inefficient and even has a bias.
The Volatility Process

Table 2.14: Mean and (Standard-Deviation) for the estimation of each parameter via EKF over $P = 500$ paths of lengths $N = 5000$ and $N = 50,000$. The true values are $(\omega^* = 0.10, \theta^* = 10, \xi^* = 0.03, \rho^* = -0.50)$.

<table>
<thead>
<tr>
<th>$N$</th>
<th>$\hat{\omega}$</th>
<th>$\hat{\theta}$</th>
<th>$\hat{\xi}$</th>
<th>$\hat{\rho}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>5000</td>
<td>0.11933899 (0.098995729)</td>
<td>11.92271488 (9.673829518)</td>
<td>0.056092146 (0.049741887)</td>
<td>-0.34321724 (0.297433861)</td>
</tr>
<tr>
<td>50000</td>
<td>0.102554592 (0.027020734)</td>
<td>10.26233092 (2.706564396)</td>
<td>0.04383931 (0.013004526)</td>
<td>-0.351998284 (0.074998408)</td>
</tr>
</tbody>
</table>

2.3.12 The Bayesian Approach

Even if our method of choice is the Classical one, it is worth going over the Bayesian philosophy and methodologies, which have some similarities but also some fundamental differences with our point of view. The MLE methodology is a Classical (Frequentist) approach where we assume that there is a set of unknown but fixed parameters. Alternatively, in the Bayesian approach the parameters are considered as random-variables with a given prior distribution. We then use the observations (the likelihood) to update these distributions and obtain the posterior distributions.

It would seem that in order to be as objective as possible and to use the observations as much as possible, one should use priors that are non-informative. However this sometimes creates degeneracy issues and one should choose a different prior for this reason.

Markov Chain Monte-Carlo’s (MCMC) include the Gibbs Sampler as well as the Metropolis-Hastings (MH) algorithm. The theoretical justification is provided by the Hammersley-Clifford theorem and the Ergodic Averaging theorem. Details could for instance be found in [34] or [163].

Briefly, the Hammersley-Clifford theorem states that having a parameter-set $\Psi$, a state $x$ and an observation $z$, we can obtain the joint distribution $p(\Psi, x, z)$ from $p(\Psi | x, z)$ and $p(x | \Psi, z)$, under some mild regularity conditions. Therefore by applying the theorem iteratively, we can break a complicated multi-dimensional estimation problem into many simple one-dimensional problems. Creating a Markov Chain $\Psi^{(i)}$ via a Monte-Carlo process, the Ergodic Averaging theorem states that the time-average of a parameter will converge towards its posterior mean.

The Gibbs Sampler

The Gibbs Sampler consists in iterative simulations from the posterior distributions.
Having a parameter-set \( \Psi = (\Psi_j)_{1 \leq j \leq J} \)

a hidden state \( x = (x_k)_{1 \leq k \leq N} \)

and an observation set \( z = (z_k)_{1 \leq k \leq N} \)

We proceed as follows:

(0) Initialize the state vector and the parameter-set to \( x^{(0)} \) and \( \Psi^{(0)} \) and choose the prior distribution \( p(\psi) \).

For each simulation index \( i \) between 1 and \( N_{\text{sims}} \) do:

(1) Simulate \( x^{(i)} \) as

\[
  x^{(i)} \sim p(x|z, \Psi^{(i-1)})
\]

(2) Simulate each parameter from its posterior conditional on partially updated parameters: For each \( j \) between 1 and \( J \)

\[
  \Psi_j^{(i)} \sim p(\psi|z, x, \Psi_0^{(i)}, ..., \Psi_{j-1}^{(i)}, \Psi_{j+1}^{(i-1)}, ..., \Psi_J^{(i-1)})
\]

with

\[
  p(\psi|z, x, ...) \propto p(z|x, \psi, ...)p(x|\psi)p(\psi)
\]

(3) Go back to step (1) and stop after \( i \) reaches \( N_{\text{sims}} \)

(4) calculate the posterior mean for each parameter after allowing a “burn-in” period

\[
  \hat{\Psi}_j = \frac{1}{N_{\text{sims}} - i_0} \sum_{i=i_0+1}^{N_{\text{sims}}} \Psi_j^{(i)}
\]

with for instance \( i_0 = N_{\text{sims}}/10 \).

It is important to note that in some cases, the prior and the posterior distributions are the same and only differ in parameters. In this case the priors are referred to as **conjugate priors**.

The justification is available for instance in [55] and could be summed up as follows: Having two random variables \( (X, Y) \) we can write

\[
  \mathbb{E}[X] = \int x p(x) dx
\]
but
\[ p(x) = \int p(x|y)p(y)dy = \int p(x|y)\int p(y|\xi)p(\xi)d\xi dy \]
therefore, we have
\[ p(x) = \int p(\xi)h(x,\xi)d\xi \]
with
\[ h(x,\xi) = \int p(x|y)p(y|\xi)dy \]
Which shows that \( p(x) \) is a stationary solution for the above integral equation, and \( h(x,\xi) \) corresponds to the limit transition density.

Similarly, it is possible to show that for a sequence \((x_k)\) generated from a Gibbs Sampler, we have
\[ P(x_k|x_0) = \int P(x_{k-1}|x_0)P(x_k, x_{k-1})dx_{k-1} \]
It is therefore possible to see that as \( k \to +\infty \) we have
\[ P(x_k|x_0) \to p(x_k) \]
and
\[ P(x_k|x_{k-1}) \to h(x_k, x_{k-1}) \]
which are the stationary marginal and transition densities.

**A Simple Illustration**

For a simple illustration consider a sequence of Normally distributed data points \( z \) with an unknown mean \( \mu \) and an unknown variance \( 1/\lambda \). The parameter \( \lambda \) is often referred to as the *precision* of the distribution.

One possible way to proceed is to choose uniform (non-informative) priors \( p(\mu) \) and \( p(\lambda) \propto 1/\lambda \) and use the known results \[34\]
\[ p(\mu|z, \sigma) = N(\bar{Z}, \sigma) \]
with \( N(m, s) \) the Normal distribution with mean \( m \) and standard-deviation \( s \) and
\[ \bar{Z} = \frac{1}{N} \sum_{k=1}^{N} z_k \]
as well as
\[ p(\lambda|z, \mu) = G\left(\frac{N}{2}, \frac{S}{2}\right) \]
with \( G(a, A) \) the previously described Gamma distribution\[34\] and
\[ S = \sum_{k=1}^{N} (z_k - \mu)^2 \]
\[34\]Note that \( G(a, A) = P(a, Ax) \) to use our previous notations.
and again

\[
\sigma = \frac{1}{\sqrt{\lambda}}
\]

We therefore know both posterior distributions and can simulate from them iteratively and perform a Gibbs sampling as described above.

For testing this, we generated a time-series of 1000 Gaussian points with a mean of \( \mu^* = 10 \) and a standard-deviation of \( \sigma^* = 5 \). We applied the Gibbs sampler via \( N_{sims} = 10,000 \) simulations and considered the average between the 1000th and 10,000th simulations. We chose initial values \( \mu_0 = 7.0 \) and \( \sigma_0 = 3.0 \) and obtained

\[
\hat{\mu} = 9.943416 \\
\hat{\sigma} = 4.816300
\]

We plotted the simulations from the posteriors in figures 2.46 and 2.47.
Figure 2.46: Gibbs Sampler for $\mu$ in $N(\mu, \sigma)$. The true value is $\mu^* = 10.0$. 
Figure 2.47: Gibbs Sampler for $\sigma$ in $N(\mu, \sigma)$. The true value is $\sigma^* = 5.0$. 
The Metropolis-Hastings Algorithm

The Gibbs Sampler is fast and simple when the posterior distributions are known and easy to sample from. However in practice, and in particular for our Stochastic Volatility problem, this often is not the case.

We assume for simplicity that we do know the posteriors for the parameters and therefore can use the Gibbs Sampler for them, however we cannot do the same for the latent state.

In this case the Metropolis-Hastings (MH) algorithm approach can be used for $x$ as follows:

(0) Initialize the state vector and the parameter-set to $x^{(0)}$ and $\Psi^{(0)}$ and choose the prior distribution $p(\psi)$. Also choose a proposal distribution $q(x|z, \Psi)$ for the state

For each simulation index between 1 and $N_{sims}$ do:

(1-a) Simulate from the proposal distribution

$$x^{(i)} \sim q(x|z, \Psi)$$

(1-b) Compare to a randomly generated uniform random-variable $u$ the ratio

$$\alpha = \min \left( 1, \frac{p(x^{(i)}|z, \Psi)q(x^{(i)}|z, \Psi)}{p(x^{(i-1)}|z, \Psi)q(x^{(i-1)}|z, \Psi)} \right)$$

and accept $x^{(i)}$ if $\alpha > u$, otherwise reject it and set $x^{(i)} = x^{(i-1)}$.

(2) Simulate $\Psi^{(i)}$ via a Gibbs Sampler.

(3) Go back to (1-a) and continue until $i$ reaches $N_{sims}$.

(4) calculate the posterior mean for each parameter after allowing a “burn-in” period

$$\hat{\Psi}_j = \frac{1}{N_{sims} - i_0} \sum_{i=i_0+1}^{N_{sims}} \Psi^{(i)}_j$$

with for instance $i_0 = N_{sims}/10$.

Two special cases are worth being mentioned:

1. First, if we simulate from the posterior, the MH ratio becomes 1.0 and every simulation will be accepted. This is therefore a Gibbs sampler.

2. Second, if we simulate from the prior, the MH ratio becomes the likelihood ratio, which makes the computation simpler. We shall use this second case...
extensively in our stochastic volatility inferences.

The justification for the MH algorithm is available for instance in [58] or [120]. The idea is to find the transition probability from $x$ to $y$ $P(x, y)$ such that for a given target density $\pi$ we would have the invariant distribution property

$$\pi(dy) = \int P(x, dy)\pi(x)dx$$

It is possible to express the transition probability $P(x, dy)$ as

$$P(x, dy) = p(x, y)dy + \left(1 - \int p(x, z)dz\right)\delta_x(dy)$$

with $\delta_x()$ the Dirac function.

The first term above corresponds to the passage probability from $x$ to a point in $dy$, and the second term to the probability of staying at $x$.

Now if the function $p(x, y)$ satisfies the reversibility condition

$$\pi(x)p(x, y) = \pi(y)p(y, x)$$

then we could see that $\pi()$ is the invariant distribution as described above.

Indeed then calling the rejection probability

$$r(x) = 1 - \int p(x, z)dz$$

we have

$$\int P(x, A)\pi(x)dx = \int \left[\int_A p(x, y)dy + r(x)\delta_x(A)\right]\pi(x)dx =$$

$$\int_A\left[\int p(x, y)\pi(x)dx\right]dy + \int_A r(x)\pi(x)dx =$$

$$\int_A\left[\int p(y, x)\pi(y)dx\right]dy + \int_A r(x)\pi(x)dx =$$

$$\int_A(1 - r(y))\pi(y)dy + \int_A r(x)\pi(x)dx =$$

$$\int_A \pi(y)dy$$

which proves that $\pi(x)$ is the invariant distribution for the transition probability $P(x, y)$.

However in practice the reversibility condition is hardly ever satisfied and therefore we need to construct an MH density that would indeed be reversible.

Taking any proposal density $q(x, y)$ we would simply write

$$p_{MH}(x, y) = q(x, y)\min\left(1, \frac{\pi(y)/q(x, y)}{\pi(x)/q(y, x)}\right)$$
Indeed then $p_{MH}(x, y)$ would be reversible and hence admit $\pi(x)$ as its invariant distribution.

**Proof:**
To see why this is, let us consider the case where $\frac{\pi(y)/q(x, y)}{\pi(x)/q(y, x)} > 1$ which means its inverse is smaller than 1. We would then have

$$p_{MH}(x, y)\pi(x) = q(x, y)\pi(x) = q(y, x)\frac{\pi(x)/q(y, x)}{\pi(x)/q(x, y)}\pi(y) = p_{MH}(y, x)\pi(y)$$

which proves the point. (QED)

One more point we need to explain is the “Blocking” technique. Having two random variables $X_1, X_2$, the product of the conditional transition densities, admits the joint distribution $\pi(X_1, X_2)$ for invariant distribution. This is why we can alternate between parameters and hidden states. Indeed

$$\int \int P_1(x_1, dy_1|x_2)P_2(x_2, dy_2|y_1)\pi(x_1, x_2)dx_1dx_2 =$$

$$\int P_2(x_2, dy_2|y_1)\left[ \int P_1(x_1, dy_1|x_2)\pi_1|x_2(x_1|x_2)dx_1 \right] \pi_2(x_2)dx_2 =$$

$$\int P_2(x_2, dy_2|y_1)\pi_1|x_2(dy_1|x_2)\pi_2(x_2)dx_2 =$$

$$\int P_2(x_2, dy_2|y_1)\pi_2|x_1(x_2|y_1)\pi_1(dy_1)dx_2 =$$

$$\pi_1(dy_1)\int P_2(x_2, dy_2|y_1)\pi_2|x_1(x_2|y_1)dx_2 =$$

$$\pi_1(dy_1)\pi_2|1(dy_2|y_1) = \pi(dy_1, dy_2)$$

which proves that $\pi(x_1, x_2)$ is the invariant distribution for this product transition probability.

**Illustration**

We use the same example as for the Gibbs sampler, only this time we simulate from the priors and use the likelihood ratio to accept or reject the simulations. We choose the priors

$$\mu \sim N(7.0, 3.0)$$

and

$$\sigma \sim \frac{1}{\sqrt{G(1/9.0, 1.0)}}$$

We obtain after $M = 10,000$ simulations

$$\hat{\mu} = 9.989504$$
\[ \hat{\sigma} = 4.797105 \]

Naturally the evolution of the Markov Chain is different from the Gibbs Sampler’s. This can be seen in figures 2.48 and 2.49.
Figure 2.48: Metropolis-Hastings algorithm for $\mu$ in $\mathcal{N}(\mu, \sigma)$. The true value is $\mu^* = 10.0$. 
Figure 2.49: Metropolis-Hastings algorithm for $\sigma$ in $N(\mu, \sigma)$. The true value is $\sigma^* = 5.0$. 
A Few Distributions

Below are a few distributions commonly used in MCMC algorithms.

The Student Cumulative Distribution Function

\[ F(x, \nu) = 1 - I \left( \frac{\nu}{\nu + x^2}, \frac{\nu}{2} \right) \]

with \( I(x, a, b) \) the Incomplete Beta Function (IBF)

\[ I(x, a, b) = \frac{B(x, a, b)}{B(1, a, b)} \]

where

\[ B(x, a, b) = \int_0^x t^{a-1}(1 - t)^{b-1} dt \]

with \( a, b \) two strictly positive parameters.

A few plots of the IBF are provided in figure 2.50.

The Inverse-Gamma (IG) Cumulative Distribution Function \( IG(a, x) \) could be defined from that of the previously defined Gamma distribution \( P(a, x) \)

\[ P(a, x) = \frac{1}{\Gamma(a)} \int_0^x e^{-t} t^{a-1} dt \]

Indeed by definition, if the random variable \( X \) is Gamma-distributed, \( Y = 1/X \) will be IG-distributed and therefore

\[ IG(a, x) = P(Y \leq x) = P(X \geq \frac{1}{x}) = 1 - P(a, \frac{1}{x}) \]

As for the densities they are related by

\[ f_{IG}(a, x) = \frac{1}{x^2} f_{G}(a, \frac{1}{x}) \]
Figure 2.50: Plots of the Incomplete Beta Function. Implementation is based on code from “Numerical Recipes in C”.
Regression Analysis

We have the following useful results as described in [34], [163] using some of the previous distributions.

Considering a univariate regression

\[ Y = \beta X + \epsilon \]

where

\[ \epsilon \sim N(0, \sigma^2) \]

We suppose we know the priors

\[ p(\beta) = N(a, A) \]

where \( a \) corresponds to the mean and \( A \) to the variance.

\[ p(\sigma^2) = IG(b, B) \]

with the density

\[ f_{IG}(x, b, B) = \frac{B^b e^{-\frac{B}{x}}}{\Gamma(b) x^{b+1}} \]

Then we have for the \( \beta \) posterior:

\[ p(\beta|Y, X, \sigma^2) \propto p(Y|X, \beta, \sigma^2)p(\beta) \propto N(a^*, A^*) \]

with

\[ a^* = \left( \frac{1}{A} + \frac{1}{2} X^t X \right)^{-1} \left( \frac{a}{A} + \frac{X^t Y}{\sigma^2} \right) \]

\[ A^* = \left( \frac{1}{A} + \frac{X^t X}{\sigma^2} \right)^{-1} \]

As for the \( \sigma^2 \) posterior we have

\[ p(\sigma^2|Y, X, \beta) \propto p(Y|X, \beta, \sigma^2)p(\sigma^2) \propto IG(b^*, B^*) \]

with

\[ b^* = T + b \]

and

\[ B^* = (Y - \beta X)^t (Y - \beta X) + B \]
Application to Gaussian SV Models (Heston)

Various MCMC approaches have been suggested for the SV problem. Jacquier, Polson and Rossi [156] were first to apply a hybrid approach between the Gibbs Sampler and the MH algorithm to a log-SV model. Kim, Shephard and Chib [169] used a slightly different approach for the same model.

We describe the method employed by Forbes, Martin and Wright (FMW) [103]. Using their notations

\[ dv_t = \kappa (\theta - v_t) dt + \sigma \sqrt{v_t} dZ_t \]

Obviously our \((\omega, \theta, \xi, \rho)\) could easily be obtained as \((\kappa \theta, \kappa, \sigma_v, \rho)\).

The algorithm becomes:

1. Initialize \(v^{(0)} = (v_k^{(0)})_{0 \leq k \leq N}\) and choose constant and therefore non-informative priors for the parameter-set\(^{35}\)

\[ \Psi = (\kappa, \theta, \sigma_v, \rho) \]

2. We simulate the state \(v_t\) from the Heston prior, we have for any time-step \(k\) between 1 and \(N\) and simulation \(i\)

\[ v_k^{(i)} = v_{k-1}^{(i)} + \kappa (\theta - v_{k-1}^{(i)}) \Delta t + \sigma \sqrt{v_{k-1}^{(i)}} \Delta t Z_{k-1} \]

As previously mentioned the MH ratio is therefore the likelihood ratio:

\[ \alpha = \min \left( 1, \frac{p(ln S|v^{(i)}, \Psi)}{p(ln S|v^{(i-1)}, \Psi)} \right) \]

where

\[ p(ln S|v, \Psi) \propto \prod_{k=1}^{N} \frac{1}{\sqrt{(1 - \rho^2) \Delta t v_{k-1}}} \exp \left\{ -\frac{1}{2(1 - \rho^2) v_{k-1} \Delta t} (\ln S_k - \mu_k)^2 \right\} \]

with

\[ \mu_k = \ln S_{k-1} + (\mu_S - \frac{1}{2} v_{k-1}) \Delta t + \frac{\rho}{\sigma_v} (v_k - [\theta \kappa \Delta t + (1 - \kappa \Delta t)v_{k-1}]) \]

Any negative variance would be rejected in the MH step.

3. The Heston equation

\[ v_k = v_{k-1} + \kappa (\theta - v_{k-1}) \Delta t + \sigma \sqrt{v_{k-1}} \Delta t Z_{k-1} \]

\(^{35}\)As before we assume for simplicity that \(\mu_S\) is known. Adding it to the parameter-set would be easy.
could be rewritten
\[
\frac{v_k - (1 - \kappa \Delta t)v_{t-1}}{\sqrt{v_{t-1} \Delta t}} = \theta \frac{\kappa \Delta t}{\sqrt{v_{t-1} \Delta t}} + \sigma_v Z_{k-1}
\]
which is a linear regression
\[
y_k = \theta x_k + e_k
\]
with
\[
y_k = \frac{v_k - (1 - \kappa \Delta t)v_{t-1}}{\sqrt{v_{t-1} \Delta t}}
\]
\[
x_k = \frac{\kappa \Delta t}{\sqrt{v_{t-1} \Delta t}}
\]
and
\[
e_k \sim N(0, \sigma_v)
\]
Hence, taking constant priors, we have
\[
\theta | \kappa, \sigma_v, v \sim N(\bar{\theta}, \sigma_{\theta})
\]
with
\[
\bar{\theta} = \frac{\sum_{k=1}^{N} x_k y_k}{\sum_{k=1}^{N} x_k^2}
\]
and
\[
\sigma_{\theta} = \sigma_v / \sqrt{\sum_{k=1}^{N} x_k^2}
\]
What is more
\[
\sigma_v^2 | \kappa, v \sim IG(N - 1, s_v^2)
\]
with
\[
s_v^2 = \sum_{k=1}^{N} (y_k - \bar{\theta} x_k)^2
\]
It is also possible to show that
\[
p(\kappa | v) \propto St(\kappa_{\hat{\kappa}}, \sigma_{\kappa}) \left( \sum_{k=1}^{N} x_k^2 \right)^{-\frac{N}{2}}
\]
where $St(m, s)$ corresponds to the Student Law of mean $m$ and standard-deviation $s$. The expressions for these mean and standard-deviations could be found in [103].
We can therefore simulate from the priors, except we have an adjustment factor 
\( \left( \sum_{k=1}^{N} x_k^2 \right)^{-\frac{1}{2}} \) to multiply the prior by. The MH ratio will therefore be

\[
\alpha = \min \left( 1.0, \frac{p(\ln S|v, \kappa(i), \theta(i), \sigma_v(i), \rho)\left(\sum_{k=1}^{N} (x_k(i))^2\right)^{-\frac{1}{2}}}{p(\ln S|v, \kappa(i-1), \theta(i-1), \sigma_v(i-1), \rho)\left(\sum_{k=1}^{N} (x_k(i-1))^2\right)^{-\frac{1}{2}}} \right)
\]

(3) As for the correlation parameter \( \rho \), we choose a Normal proposal distribution and use a constant prior again. Therefore

\[
\alpha = \min \left( 1.0, \frac{p(\ln S|v, \kappa, \theta, \sigma_v, \rho(i))}{p(\ln S|v, \kappa, \theta, \sigma_v, \rho(i-1))} \right) \left( \frac{q(\rho(i)|v, \kappa, \theta, \sigma_v, S)}{q(\rho(i-1)|v, \kappa, \theta, \sigma_v, S)} \right)
\]

with \( q() \) the Normal distribution with mean

\[
\frac{\sum_{k=1}^{N} x_k y_k}{\sum_{k=1}^{N} x_k^2}
\]

and variance

\[
\frac{\Delta t}{\sum_{k=1}^{N} x_k^2}
\]

with

\[
x_k = \frac{v_k - \kappa \theta \Delta t - (1 - \kappa \Delta t) v_{k-1}}{\sigma_v \sqrt{\theta_{k-1}}}
\]

\[
y_k = \frac{\ln S_k - \ln S_{k-1} - (\mu_S - \frac{1}{2} v_{k-1}) \Delta t}{v_{k-1}}
\]

Note that for any of the parameters above if we simulate one that does not satisfy the usual constraints \( \theta \geq 0, \kappa \geq 0, \sigma_v \geq 0, \sigma_s \leq 2\kappa \theta \) and \(-1 \leq \rho \leq 1\), the we simply do not accept them during the MH accept/reject step.

Also note that we update \( (\kappa, \theta, \sigma_v) \) in a “block” instead of updating them one by one. This technique is used by many since it makes the algorithm faster.

For the actual results the reader could refer to Forbes et al. [103]. Note that the authors test their Bayesian Estimator against simulated data, and observe inefficiency. This is in agreement with our observations when applying MLE techniques to simulated data.
2.3.13 Using the Characteristic Function

In a recent article [31] the use of the Characteristic Function has been suggested for the purpose of Filtering. In this approach however, we have to limit ourselves to the case where
\[ F(U, V, x_t) = \mathbb{E}[\exp(U z_{t+1} + V x_{t+1})|x_t] = \exp\{C(U, V) + D(U, V)x_t\} \]

One natural form would be the affine process where
\[ F(U, V, x_t) = \mathbb{E}[\exp(U z_{t+1} + V x_{t+1})|x_t] = \exp\{C(U, V) + D(U, V)x_t\} \]

After choosing the initial conditions, the Time Update equation
\[ p(z_{t+1}, x_{t+1}|t) = \int p(z_{t+1}, x_{t+1}|x_t)p(x_t|t)dx_t \]

becomes in terms of the characteristic function
\[ G_{t+1}(V) = \frac{\int_{-\infty}^{+\infty} F_{z_{t+1}x_t}(iU, V) \exp(-iU z_{t+1})dU}{\int_{-\infty}^{+\infty} F_{x_t}(iU, 0) \exp(-iU z_{t+1})dU} \]

This remarkably gives us a one-step induction expression
\[ G_{t+1}|t+1(V) = \frac{\int_{-\infty}^{+\infty} \exp[C(iU, V) - iU z_{t+1}]G_{t+1}(D(iU, V))dU}{\int_{-\infty}^{+\infty} \exp[C(iU, 0) - iU z_{t+1}]G_{t+1}(D(iU, 0))dU} \]

which allows us to determine the a posteriori estimate and errors
\[ \hat{x}_t = G_{t+1}(0) \]

and
\[ P_t = \text{Var}(x_t) = G_{t+1}''(0) - \left(G_{t+1}'(0)\right)^2 \]

at each iteration.

In this framework, the likelihood function could be written as
\[ L_{1:T} = \prod_{t=0}^{T-1} l_t \]
with
\[ l_t = \frac{1}{2\pi} \int_{-\infty}^{+\infty} \exp[C(iU, 0) - iU z_{t+1}] G_{t|t}[D(iU, 0)] dU \]
which is equivalent to
\[ l_t = \frac{1}{\pi} \int_{0}^{+\infty} \Re \{ \exp[C(iU, 0) - iU z_{t+1}] G_{t|t}[D(iU, 0)] \} dU \tag{2.30} \]
where \( \Re \{ \} \) corresponds to the real part of a complex number.

In order to be able to calculate the integral, we need to know the value of \( G_{t|t}(x) \) at each point. For this, Bates [31] suggests making an assumption on the distribution of the hidden state. For a Gamma distribution we have a moment-generating-function of the form
\[ G_{t|t}(x) = (1 - \kappa x)^{-\nu_t}. \]

The integral (2.30) can be evaluated numerically, however when dealing with “outliers” the density of the observation takes near-zero values which makes the integration difficult. Bates suggests scaling transformations equivalent to the importance sampling technique used in particle filtering.

Independently from this, Dragulescu and Yakovenko [81], [219] derived a semi-analytic expression for the likelihood under the Heston model, by using Fourier Inversion. Note that a Particle Filter calculates this very integral via Monte-Carlo simulations.

It is worth noting that the main advantage of our Particle Filtering approach is its complete generality. Indeed the Bates method would work only for model classes that have an exponentially affine Fourier Transform. It is true that the Heston model falls in this category, however a VGG (Variance Gamma with Gamma-distributed Arrival-Rate) process would not, and therefore could only be analyzed though a simulation-based methodology.
2.3.14 Introducing Jumps

The Model

As in Bates [28], let us introduce a Jump Process (independent from the Brownian Motion) with a given intensity $\lambda$ and a fixed $j$ fractional jump size $0 \leq j < 1$. The number of jumps between $t$ and $t + dt$ will therefore be $dN_t$. Needless to say, if either the intensity $\lambda = 0$ or the jump-size $j = 0$, then we are back to the pure diffusion case.

The new Stochastic Differential Equation for the Stock Price in the risk-neutral framework will be

$$dS_t = (\mu_S + \lambda j)S_t dt + \sqrt{v_t}S_t dB_t - S_t j dN_t$$

and applying Ito’s lemma for Semi-Martingales

$$d \ln S_t = (\mu_S - \frac{1}{2}v_t + \lambda j)dt + \sqrt{v_t}dB_t + \ln(1 - j)dN_t$$

which we can rewrite in the discrete version as

$$\ln S_{k+1} = \ln S_k + (\mu_S - \frac{1}{2}v_k + \lambda j)\Delta t + \sqrt{v_k}\sqrt{\Delta t}B_k + \mu_k$$

with $\mu_0 = 0$ and

$$\mu_k = \delta_0(0)e^{-\lambda \Delta t} + \delta_0(\ln(1 - j)) (1 - e^{-\lambda \Delta t})$$

where $\delta_0()$ corresponds to the Dirac delta function.\footnote{37}

Also

$$v_k = v_{k-1} + (\omega - \theta v_{k-1})\Delta t + \xi \sqrt{v_{k-1}}\sqrt{\Delta t}Z_{k-1} - \rho \xi [\ln S_{k-1} + (\mu_S + \lambda j - \frac{1}{2}v_{k-1})\Delta t + \sqrt{v_{k-1}}\sqrt{\Delta t}B_{k-1} + \mu_{k-1} - \ln S_k]$$

which completes our set of equations.

It is important to note that the new parameter-set is

$$\Psi = (\omega, \theta, \xi, \rho, \lambda, j)$$

which effectively gives us two additional degrees of freedom.\footnote{38}

\footnote{36}We could make $j$ a Gaussian random variable without changing the methodology.\footnote{37}This means that $-\infty < \mu_k \leq 0$ for every $k$.\footnote{38}Note that we are assuming that we can have at most one jump within $[t, t + \Delta t]$ which means that $\Delta t$ is small enough. This is completely different from pure-jump models such as Variance-Gamma.\footnote{38}A related idea was developed by Hamilton [126] as well as Chourdakis [59] and Deng [72]. Chourdakis uses the characteristic function for the jump-diffusion process. Doucet [80] suggests the use of Particle Filtering for the Jump process. Maheu & McCurdy [184] use a fully integrated GARCH likelihood with Poisson Jumps. Ait-Sahalia [3] uses moments to separate the diffusion parameters from the jumps. Johannes, Polson and Stroud [164] use the Particle Filtering technique as well, however in a Bayesian MCMC framework. Finally Honoré [142] shows that an MLE approach always works for a constant jump size.
The Generic Particle Filter

Since $\mu_k$ is following a Poisson process we have to use a Non-Gaussian Filter. The use of a Generic Particle Filter (GPF) is therefore natural. In a generic particle filter the proposal distribution $q(x_k)$ is simply set equal to $p(x_k|x_{k-1})$.

The state $x_k$ could be chosen as

$$x_k = \begin{pmatrix} \mu_k \\ v_k \end{pmatrix}$$

and the transition equation becomes

$$x_k = \begin{pmatrix} \\
\delta_0(0)e^{-\lambda \Delta t} + \delta_0(\ln(1-j))(1 - e^{-\lambda \Delta t}) \\
v_{k-1} + [(\omega - \rho \xi (\mu_S + \lambda j) - (\theta - \frac{1}{2} \rho \xi) v_{k-1}] \Delta t + \rho \xi [\ln(\frac{S_k}{S_{k-1}}) - \mu_{k-1}] + \xi \sqrt{1 - \rho^2} \sqrt{v_{k-1} \Delta Z_{k-1}} \end{pmatrix}$$

It becomes therefore possible to implement a Particle Filter as follows

1. Choose $v_0$ and $P_0 > 0$ and set $\mu_0 = 0$, so for $i$ in $1, ..., N_{sims}$

$$x_0^{(i)} = \begin{pmatrix} 0 \\ v_0 + \sqrt{P_0} Z^{(i)} \end{pmatrix}$$

Then for each $k$ with $1 \leq k \leq N$ do

2. Write the new $\tilde{x}_k^{(i)} = (\tilde{\mu}_k^{(i)}, \tilde{v}_k^{(i)})^t$ as the result of simulations

$$\tilde{v}_k^{(i)} \sim \mathcal{N} \left( m = v_{k-1}^{(i)} + [\omega - \rho \xi (\mu_S + \lambda j) - (\theta - \frac{1}{2} \rho \xi) v_{k-1}^{(i)}] \Delta t + \rho \xi [\ln(\frac{S_k}{S_{k-1}}) - \mu_{k-1}^{(i)}], s \right)$$

with $s = \xi \sqrt{1 - \rho^2} \sqrt{v_{k-1}^{(i)} \Delta t}$

and

$$\tilde{\mu}_k^{(i)} = 0$$

if $\mathcal{U}[0,1] \leq e^{-\lambda \Delta t}$ and

$$\tilde{\mu}_k^{(i)} = \ln(1 - j)$$

otherwise.

3. Define the weights

$$w_k^{(i)} = w_{k-1}^{(i)} p(z_k|\tilde{x}_k^{(i)})$$

with

$$p(z_k|\tilde{x}_k^{(i)}) = n(z_k, z_{k-1} + (\mu_S + \lambda j - \frac{1}{2} \tilde{v}_k^{(i)}) \Delta t + \tilde{\mu}_k^{(i)}, \sqrt{\tilde{v}_k^{(i)} \Delta t})$$
(4) Normalize the weights
\[ \tilde{w}_k^{(i)} = \frac{w_k^{(i)}}{\sum_{i=1}^{N_{sims}} w_k^{(i)}} \]

(5) Resample the points \( \tilde{x}_k^{(i)} \) and get \( x_k^{(i)} \) and reset \( w_k^{(i)} = \tilde{w}_k^{(i)} = 1/N_{sims} \).

This completes the Generic Particle Filtering algorithm.

Note that there is no Kalman Filtering here and therefore
\[ \hat{\tilde{z}}_k = \frac{1}{N_{sims}} \sum_{i=1}^{N_{sims}} \tilde{z}_k^{(i)} \]

with \( \tilde{z}_k^{(i)} \) the estimation of \( z_k \) from \( \tilde{x}_{k-1}^{(i)} \)
\[ \tilde{z}_k^{(i)} = z_{k-1} + (\mu_S + \lambda j - \frac{1}{2} v_{k-1}^{(i)}) \Delta t + \mu_{k-1} \]
and the estimation error is \( z_k - \hat{\tilde{z}}_k \) as before.

The likelihood maximization is not different from the EPF or UPF. We need to maximize
\[ \sum_{k=1}^{N} \ln \left( \frac{1}{N_{sims}} \sum_{i=1}^{N_{sims}} w_k^{(i)} \right) \]

where \( w_k^{(i)} \)'s are defined at Step 3.

**Extended/ Unscented Particle Filters**

Using the same model, we can take advantage of the independence of \( v_k \) and \( \mu_k \) and apply the (Nonlinear) Gaussian Kalman Filter to the former.

In this case, the Steps 2 and 3 should be replaced with:

(2-a) Write \( \hat{x}_k^{(i)} = (\hat{\mu}_k^{(i)}, \hat{v}_k^{(i)})^t \) with
\[ \hat{v}_k^{(i)} = KF(v_{k-1}^{(i)}) \]
with \( P_k^{(i)} \) the associated a posteriori error covariance matrix, KF the Extended or Unscented Kalman Filter, and
\[ \hat{\mu}_k^{(i)} = \mu_{k-1} \]

(2-b) Now take the simulations
\[ \tilde{v}_k^{(i)} \sim N \left( \hat{v}_k^{(i)}, P_k^{(i)} \right) \]
and

\[ \hat{\mu}_k^{(i)} = 0 \]

if \( U[0, 1] \leq e^{-\lambda \Delta t} \) and

\[ \hat{\mu}_k^{(i)} = \ln(1 - j) \]

otherwise.

(3) Define the weights

\[ w_k^{(i)} = w_{k-1}^{(i)} \frac{p(z_k | \tilde{x}_k^{(i)} p(\tilde{x}_k^{(i)} | x_{k-1}^{(i)})}{q(\tilde{x}_k^{(i)} | x_{k-1}^{(i)}, z_{1:k})} \]

with

\[ p(z_k | \tilde{x}_k^{(i)}) = n(z_k, z_{k-1} + (\mu_S + \lambda j - \frac{1}{2} \tilde{v}_k^{(i)}) \Delta t + \tilde{\mu}_k^{(i)}, \sqrt{\tilde{v}_k^{(i)} \Delta t}) \]

\[ p(\tilde{x}_k^{(i)} | x_{k-1}^{(i)}) = n(\tilde{v}_k^{(i)}, \mathbf{m}, \mathbf{s} = \xi \sqrt{1 - \rho^2} \sqrt{\frac{1}{v_{k-1}^{(i)} \Delta t} \tilde{v}_k^{(i)} \Delta t} \tilde{\mu}_k^{(i)} \mu_{k-1}^{(i)}) \]

with

\[ \mathbf{m} = v_{k-1}^{(i)} + [\omega - \rho \xi (\mu_S + \lambda j) - (\theta - \frac{1}{2} \rho \xi) v_{k-1}^{(i)}] \Delta t + \rho \xi \ln\left(\frac{S_k}{S_{k-1}}\right) - \rho \xi \mu_{k-1}^{(i)} \]

and

\[ q(\tilde{x}_k^{(i)} | x_{k-1}^{(i)}, z_{1:k}) = n(\tilde{v}_k^{(i)}, \tilde{v}_k^{(i)}), \mathbf{P}_k^{(i)} p(\tilde{\mu}_k^{(i)} | \mu_{k-1}^{(i)}) \]

Note that as for the GPF, the terms \( p(\tilde{\mu}_k^{(i)} | \mu_{k-1}^{(i)}) \) cancel out and need not to be evaluated.

The rest of the algorithm remains the same.
This way we will not lose the information contained in the Kalman gain for the Gaussian dimension.

**The Srivastava Approach**

Srivastava [222] suggests the following approach for simulating the jump component:
Instead of allowing a jump at each time-interval \([t_k, t_k + \Delta t]\) with a probability \( 1 - e^{-\lambda \Delta t} \) as we do now, we can flag the time-steps such that

\[ t_{k-1} < \frac{1}{\lambda} \ln \left( \frac{1}{U[0, 1]} \right) \leq t_k \]

where \( U[0, 1] \) is a uniform random-variable between zero and one, and then perform a jump of size \( |\ln(1 - j)| \) on these steps for all paths.
We therefore would first initialize $t_p = 0$ and loop through $k$’s between 1 and $N$ and if
\[ e^{-\lambda(t_k - t_p)} \leq U[0, 1] < e^{-\lambda(t_{k-1} - t_p)} \]
we flag this $k$ and set $t_p = t_k$ and resimulate $U[0, 1]$.

In the Particle Filter we would set for all indices $i$’s
\[ \tilde{\mu}_k^{(i)} = \ln(1 - j) \]
for the flagged $k$’s and we would set $\tilde{\mu}_k^{(i)} = 0$ for other indices.

It is important to note that in this approach the simulation for the Jump component is completely “orthogonal” to the Diffusion SIS part. Indeed the index $i$ above is irrelevant for the entity $\tilde{\mu}_k^{(i)}$. This means that the KF step, the weight calculation and the resampling are independent from the Jump component altogether.

**Numeric results**

As a check, we simulate a time-series with the parameter-set
\[ \Psi^* = (\omega^* = 0, \theta^* = 0, \xi^* = 0, \rho^* = 0, \lambda^* = 2.52, j^* = 0.20) \]
which corresponds to a jump frequency of $\lambda \Delta t = 0.01$ and a jump-size of 20 percent. We generated $N = 245$ points and used $M = 1000$ particles.

The estimated set via the above EPF is
\[ \hat{\Psi} = (\hat{\omega} = 0.23, \hat{\theta} = 1.5, \hat{\xi} = 0.34, \hat{\rho} = 0.21, \hat{\lambda} = 2.65, \hat{j} = 0.20) \]

As we see the diffusion parameters are not close to the original ones, but this is probably due to the small $\Delta t$ as previously discussed. The jump parameters are close to the original ones, which means the filter is valid for the jump component. Note that despite the difference in the diffusion parameters, the estimated and original time-series are rather close for a new simulation, as could be seen in figure 2.52.

This reconfirms our previous remark: when the parameters affect the drift of the observation (as opposed to its noise) their estimation is far more accurate and requires fewer data points.

**The Optimization Algorithm**

It is important to realize that the Likelihood function here (due to the jumps) is not Differentiable everywhere and therefore gradient based maximization methods could not be applied. The optimization could however still be carried out via the Direction Set algorithm as previously described.

Note that as mentioned in [164] so far there has been no formal proof on the
convergence of the discretized Jump-Diffusion equations towards the continuous ones, however empirical evidence makes the convergence assumption plausible.
Figure 2.51: Comparison of EPF results for Heston and Heston+Jumps models. The presence of jumps can be seen in the residuals.
Figure 2.52: Comparison of EPF results for Simulated and Estimated Jump-Diffusion Time-Series. The filtered data matches the real data fairly well.
2.3.15 Pure-Jump Models

The Variance Gamma with Stochastic Arrival (VGSA) and the Variance Gamma with Gamma Arrival (VGG) Models were defined in the previous chapter. These models being non-Gaussian, we could apply the Particle Filtering technique to them.

It is important to note that we are not dealing with Diffusion models and therefore we do not have the Girsanov theorem. We are estimating the parameter-set

\[ \Psi = (\mu_S, \theta, \sigma, \nu, ...) \]

In order to make the back-testing simpler we suppose however that we know the stock-drift and try to estimate the other parameters. However as mentioned earlier, for a high frequency data-set we have

\[ \Delta t = o(\sqrt{\Delta t}) \]

and the drift term has a negligible impact.

VG

The Variance Gamma model has the advantage of offering an integrated density, which allows us to calculate the exact likelihood.

Calling \( z = \ln(S_k/S_{k-1}) \) and \( h = t_k - t_{k-1} \) and posing \( x_h = z - \mu_S h - \frac{\theta}{\nu} \ln(1 - \theta \nu - \sigma^2 \nu/2) \) we have

\[
p(z|h) = \frac{2 \exp(\theta x_h/\sigma^2)}{\nu^{\frac{\nu}{2}} \sqrt{2\pi \sigma \Gamma(\frac{\nu}{2})}} \left( \frac{x_h^2}{2\sigma^2/\nu + \theta^2} \right)^{\frac{\nu}{2} - \frac{1}{2}} K^{\frac{\nu}{2} - \frac{1}{2}} \left( \frac{1}{\sigma^2} \sqrt{\frac{x_h^2 (2\sigma^2/\nu + \theta^2)}{2\nu/\sigma^2}} \right)
\]

and the likelihood is

\[ L_{1:N} = \prod_{k=1}^{N} p(z_k|z_{k-1}, h) \]

The implementation of the above estimation procedure is straightforward and has already been done in [182].

One could also back-test the estimation procedure in the following way: First choose a parameter-set \( (\theta, \sigma, \nu) \) as well as a drift \( \mu_S \) and a time-step \( \Delta t \). Then simulate via Monte-Carlo a Gamma-distributed random-variable as well as a Gaussian one. Deduce an artificial stock-price time-series and apply the MLE procedure to it and try to recover the original parameter-set.

Using \( \Delta t = 1/252, \mu^*_S = 0.05 \) and

\[
\begin{align*}
\theta^* &= 0.02 \\
\sigma^* &= 0.2 \\
\nu^* &= 0.005
\end{align*}
\]
We simulated 500 data points, applied the MLE and found an optimal parameter-set \( \hat{\Psi} = (0.018, 0.22, 0.006) \) which is close to the original set.

**VGSA**

Using the same notations as in the previous chapter, the Euler discretized VGSA process could be written via the auxiliary variable
\[
y_k = y_{k-1} + \kappa(\eta - y_{k-1})\Delta t + \lambda\sqrt{y_{k-1}}\Delta t W_{k-1}
\]
and the state
\[
x_k = F_{\nu^{-1}}(y_k \Delta t, U[0, 1])
\]
as well as the observation
\[
z_k = \ln S_{k+1} + z_{k-1} + (\mu_S + \omega) \Delta t + \theta x_k + \sigma \sqrt{x_k} B_k
\]
with \( \omega = \frac{1}{\nu} \ln(1 - \theta \nu - \sigma^2 \nu/2) \).

**The Filtering Algorithm**

The PF algorithm could therefore be written as follows:

1. **Initialize** the arrival-rate \( y_0^{(j)} \), the state \( x_0^{(i)} \) and the weight \( w_0^{(i)} \) for \( j \) between 1 and \( M_{sims} \), and \( i \) between 1 and \( N_{sims} \)

While \( 1 \leq k \leq N \)

2. **Simulate** the arrival-rate \( y_k^{(j)} \) for \( j \) between 1 and \( M_{sims} \)

\[
y_k^{(j)} = y_{k-1}^{(j)} + \kappa(\eta - y_{k-1}^{(j)})\Delta t + \lambda\sqrt{y_{k-1}^{(j)}}\Delta t N^{-1} \left( U^{(j)}[0, 1] \right)
\]

3-a. **Simulate** the state \( x_k^{(i)} \) for each \( y_k^{(j)} \) and for \( i \) between 1 and \( N_{sims} \)

\[
\hat{x}_k^{(i)} = F_{\nu^{-1}}(y_k^{(j)} \Delta t, U^{(i)}[0, 1])
\]

3-b. **Compute** the unconditional state

\[
\hat{x}_k^{(i)} = \int \hat{x}_k^{(i)}(y_k)p(y_k|y_{k-1})dy_k \approx \frac{1}{M_{sims}} \sum_{j=1}^{M_{sims}} \hat{x}_k^{(i,j)}
\]

4. **Calculate** the associated **weights** for each \( i \)

\[
w_k^{(i)} = w_{k-1}^{(i)}p(z_k|\hat{x}_k^{(i)})
\]
with
\[ p(z_k|x_k^{(i)}) = n(z_k, m, s) \]
the normal density with mean
\[ m = z_{k-1} + (\mu_S + \omega) \Delta t + \theta \bar{x}_k^{(i)} \]
and standard deviation
\[ s = \sigma \sqrt{x_k^{(i)}} \]

(5) Normalize the weights
\[ \tilde{w}_k^{(i)} = \frac{w_k^{(i)}}{\sum_{i=1}^{N_{sims}} w_k^{(i)}} \]

(6) Resample the points \( \bar{x}_k^{(i)} \) and get \( x_k^{(i)} \) and reset \( w_k^{(i)} = \tilde{w}_k^{(i)} = 1/N_{sims} \).

(7) Increment \( k \), Go back to step (2) and Stop at the end of the While loop.
Parameter Estimation

As usual, the Log-Likelihood to be maximized is

\[ \ln(L_{1:N}) = \sum_{k=1}^{N} \ln \left( \sum_{i=1}^{N_{sims}} w_k^{(i)} \right) \]

The maximization takes place over the parameter-set \( \Psi = (\kappa, \eta, \lambda, \nu, \theta, \sigma) \). Again, in reality the stock-drift \( \mu \) should be estimated together with the other parameters, however with a view to simplifying, we suppose we know \( \mu \) in our back-testing procedures.

A More Efficient Algorithm

We could take advantage of the fact that VG provides an integrated density of stock return. Calling \( z = \ln(S_k/S_{k-1}) \) and \( h = t_k - t_{k-1} \) and posing \( x_h = z - \mu_h h - \frac{h}{\nu} \ln(1 - \theta \nu - \sigma^2 \nu/2) \) we have

\[ p(z|h) = \frac{2 \exp(\theta x_h/\sigma^2)}{\nu^{\frac{h}{\nu} \sqrt{2\pi \sigma^2}}} \left( \frac{x_h^2}{2\sigma^2/\nu + \theta^2} \right)^{\frac{h}{\nu} - \frac{1}{2}} K_{\frac{h}{\nu} - \frac{1}{2}} \left( \frac{1}{\sigma^2 \sqrt{2\pi \sigma^2}} \right) \]

As we can see the dependence on the Gamma distribution is “integrated out” in the above.

For VGSA, for a given arrival rate \( dt^* = y \) we have a VG distribution and

\[ d \ln S_t = (\mu + \omega) dt + B(\gamma(dt^*, 1, \nu); \theta, \sigma) \]

and the corresponding integrated density becomes

\[ p(z|h, h^*) = \int_0^{+\infty} p(z|g, h) p(g|h) dg \]

with \( p(z|g, h) \) a Normal density and \( p(g|h) \) a Gamma density.

More accurately

\[ p(z|g, h) = \frac{1}{\sigma \sqrt{2\pi g}} \exp \left( -\frac{1}{2\sigma^2 g} \left( z - \mu \frac{h}{\nu} - \frac{h}{\nu} \ln(1 - \theta \nu - \sigma^2 \nu/2) - \theta g \right)^2 \right) \]
and
\[ p(g|h) = \frac{g^{\frac{1}{\nu}} \exp\left(-\frac{g}{\nu}\right)}{\nu^{\frac{1}{\nu}} \Gamma\left(\frac{1}{\nu}\right)} \]

Now, for VGSA we simply have a different arrival rate \( h^* \) for the Gamma process and therefore
\[ p(z|h, h^*) = \int_0^{+\infty} p(z|g, h)p(g|h^*)dg \]
which demonstrates the point.

This gives us the idea of using the arrival rate as the state and use the following algorithm:

1. **Initialize** the state \( x_0^{(i)} \) and the weight \( w_0^{(i)} \) for \( i \) between 1 and \( N_{sims} \)

2. **Simulate** the state \( x_k^{(i)} \) for \( i \) between 1 and \( N_{sims} \)

   \[ \tilde{x}_k^{(i)} = x_{k-1}^{(i)} + \kappa(\eta - x_{k-1}^{(i)})\Delta t + \lambda \sqrt{x_{k-1}^{(i)}} \sqrt{\Delta t} \mathcal{N}^{-1}\left(\mathcal{U}(i)[0, 1]\right) \]

3. **Calculate** the associated weights for each \( i \)

   \[ w_k^{(i)} = w_{k-1}^{(i)} p(z_k|\tilde{x}_k^{(i)}) \]

   with \( p(z_k|\tilde{x}_k^{(i)}) \) as defined in (2.31) where \( h \) will be set to \( \Delta t \) and \( h^* \) to the simulated state \( \tilde{x}_k^{(i)} \) times \( \Delta t \)

4. **Normalize** the weights

   \[ \tilde{w}_k^{(i)} = \frac{w_k^{(i)}}{\sum_{i=1}^{N_{sims}} w_k^{(i)}} \]

5. **Resample** the points \( \tilde{x}_k^{(i)} \) and get \( x_k^{(i)} \) and reset \( w_k^{(i)} = \tilde{w}_k^{(i)} = 1/N_{sims} \).

6. **Increment** \( k \), Go back to step (2) and **Stop** at the end of the **While** loop.

The advantage of this method is that there is one simulation process instead of two and we “skip” the Gamma distribution altogether. However the dependence of the observation \( z_k \) on \( x_k \) is highly nonlinear which makes the convergence more difficult.

**An Extended/ Unscented Particle Filter**

Finally, a natural idea would be to use a proposal distribution \( q(x) \) for the state taking into account the observation information. In order to be able to
use a Kalman-based proposal distribution (EPF or UPF) we need a Gaussian approximation. Note that given the strong non-Gaussianity of the equations, we absolutely need the Particle Filtering aspect. The Gaussian approximation for the observation equation would be

\[ z_k = z_{k-1} + (\mu_S + \omega \theta x_k) \Delta t + \sqrt{\theta^2 \nu + \sigma^2} \sqrt{x_k \Delta t} B_k \]

which is of the form \( z_k = h(x_k, B_k) \) and allows us to use the Kalman Filtering algorithm.

We therefore replace the steps (2) and (3) of the previous algorithm with:

**2-a** Apply an Extended/Unscented Kalman Filter for \( i \) between 1 and \( N_{sims} \) to the state \( x^{(i)}_{k-1} \) and obtain

\[ \hat{x}^{(i)}_k = KF(x^{(i)}_{k-1}) \]

as well as the associated covariance \( P^{(i)}_k \).

**2-b** Simulate the state for \( i \) between 1 and \( N_{sims} \)

\[ \tilde{x}^{(i)}_k = \hat{x}^{(i)}_k + \sqrt{P^{(i)}_k} \mathcal{N}^{-1} (U^{(i)}[0,1]) \]

**3** Calculate the associated weights for each \( i \)

\[ w^{(i)}_k = w^{(i)}_{k-1} \frac{p(z_k|\tilde{x}^{(i)}_k)p(\tilde{x}^{(i)}_k | x^{(i)}_{k-1})}{q(\tilde{x}^{(i)}_k | x^{(i)}_{k-1}, z_{1:k})} \]

with \( p(z_k|\tilde{x}^{(i)}_k) \) as defined in (2.31) where \( h \) will be set to \( \Delta t \) and \( h^* \) to the simulated state \( \tilde{x}^{(i)}_k \) times \( \Delta t \)

\( p(\tilde{x}^{(i)}_k | x^{(i)}_{k-1}) \) the Normal density with mean \( x^{(i)}_{k-1} + \kappa (\eta - x^{(i)}_{k-1}) \Delta t \) and standard deviation \( \lambda \sqrt{x^{(i)}_{k-1}} \sqrt{\Delta t} \)

\( q(\tilde{x}^{(i)}_k | x^{(i)}_{k-1}, z_{1:k}) \) the Normal density with mean \( \hat{x}^{(i)}_k \) and standard deviation \( \sqrt{P^{(i)}_k} \).

The rest of the algorithm is exactly the same as the previous one.

**Numeric Results**

We performed the same kind of back-testing procedure as for the VG model, using either the above Particle-Filter to an artificially generated stock-price time-series. We chose \( \Delta t = 1/252, \mu_S^* = 0, \) \( y_0 = 1 \) and

\[ \Psi^* = (\kappa^* = 0, \eta^* = 0, \lambda^* = 0, \nu^* = 0.005, \theta^* = 0.02, \sigma^* = 0.2) \]

\(^{39}\)We are using the fact that for \( X(t) = B(\gamma(t, 1, \nu); \theta, \sigma) \) we have a mean \( \theta t \) and a variance \( (\theta^2 \nu + \sigma^2) t \) as stated in [182].
after applying the Importance Sampling / Resampling PF via the modified Bessel function, we found
\[
\hat{\Psi} = (0.13, 0.001, 0.37, 0.0048, 0.018, 0.21)
\]

which seems to indicate that the estimation process for \((\nu, \theta, \sigma)\) works well, while the one for \((\kappa, \eta, \lambda)\) does not. However if we simulate two sets of spot-price timeseries with these different parameter-sets we will see that the generated paths are very similar. See Figures 2.53 and 2.54. This also confirms our previous remarks about the estimation of the parameters affecting the noise.

We performed a second test with more realistic choice of parameters. With once again \(\Delta t = 1/252, N_{\text{sims}} = 100\) and 500 data points corresponding to two years.

The real values were
\[
\Psi^* = (\kappa^* = 2.10, \eta^* = 5.70, \lambda^* = 2.00, \nu^* = 0.05, \theta^* = -0.40, \sigma^* = 0.20)
\]

Note that \(\theta\) has a negative value which corresponds to the negative skewness of the distribution.

We choose a fairly reasonable initial set
\[
\Psi_0 = (\kappa_0 = 2.00, \eta_0 = 6.00, \lambda_0 = 1.50, \nu_0 = 0.03, \theta_0 = -0.30, \sigma_0 = 0.30)
\]

and
\[
\mu_0 = \mu^* = 0.05
\]

We find the optimal parameter-set
\[
\hat{\Psi} = (\hat{\kappa} = 4.25, \hat{\eta} = 7.89, \hat{\lambda} = 3.25, \hat{\nu} = 0.047, \hat{\theta} = -0.40, \hat{\sigma} = 0.19)
\]

and
\[
\hat{\mu} = \mu^* = 0.05
\]

Again we see that the estimation for the three VG parameters \((\nu, \theta, \sigma)\) is much more accurate than those corresponding to the arrival process \((\kappa, \eta, \lambda)\). This, despite our choosing the initial arrival parameters close to the real ones. As previously stated, the time-series of spot-prices has little sensitivity to the arrival-rate parameters and a higher degree of sensitivity to the Gamma process parameters. Again, this shows that Estimation methodologies such as MLE work much better when applied to parameters that affect the drift of an observation, and not its noise.
Figure 2.53: The simulated arrival rates via $\Psi = (\kappa = 0, \eta = 0, \lambda = 0, \sigma = 0.2, \theta = 0.02, \nu = 0.005)$ and $\Psi = (\kappa = 0.13, \eta = 0, \lambda = 0.40, \sigma = 0.2, \theta = 0.02, \nu = 0.005)$ are quite different ...
Figure 2.54: ... However the simulated log stock Prices are close.
Diagnostics

As for diagnostics, we need to estimate the observation error associated with the algorithm. We define once again

$$\hat{z}_k^{(i)} = z_{k-1} + (\mu_S + \omega + \theta \tilde{x}_k^{(i)}) \Delta t$$

or

$$\hat{z}_k^- = \frac{1}{N_{sims}} \sum_{i=1}^{N_{sims}} \hat{z}_k^{(i)}$$

and the error term

$$e_k = z_k - \hat{z}_k^-$$

The variance associated to this error is

$$s_k = (\theta^2 \nu + \sigma^2) \frac{1}{N_{sims}} \sum_{i=1}^{N_{sims}} \tilde{x}_k^{(i)} \Delta t$$

and

$$\hat{e}_k = e_k / s_k$$

would represent the normalized error.

MPE/ RMSE

In order to measure the performance, once again we use the Mean Price Error (MPE) and the Root Mean-Squared Error (RMSE). As an example we use the S&P500 data between 1992 and 1994 (as used in [182]) we find for the generic particle filter (GPF) and the extended particle filter (EPF)

<table>
<thead>
<tr>
<th></th>
<th>MPE</th>
<th>RMSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>PF</td>
<td>-0.000350241</td>
<td>0.005867065</td>
</tr>
<tr>
<td>EPF</td>
<td>-4.74747e-07</td>
<td>0.005869782</td>
</tr>
</tbody>
</table>

Table 2.15: MPE and RMSE for the VGSA model under a generic PF as well as the EPF.

As we can see the use of the extended Kalman filter as the proposal distribution brings some improvement.

Chi-Square Test

The residuals are normal, indeed a $\chi^2_{20}$ test provides us with a value of 10.397699 which is below the threshold value of 31.5 for a confidence of 0.95. This means that the non-Gaussianity was “filtered out” of the time-series successfully. This
could also be observed in the corresponding histogram in figure 2.57.

**Auto-Correlation**
Having $p = 7$ parameters and taking $K = 27$, we shall have $K - p = 20$ so we will compare the output of the Box-Ljung test to the $\chi^2_{20}$ threshold, which as previously mentioned for a confidence of 0.95 is around 31.5. We find a value of 0.001138 which definitely passes the test. This shows that the residuals are indeed uncorrelated.

**Variogram**
The variogram still indicates that we have independent and identically distributed random variables. Calling

$$
\gamma_h = \frac{1}{2} \mathbb{E}[(\tilde{e}_{k+h} - \tilde{e}_k)^2] = \frac{1}{2} \mathbb{E}[\tilde{e}_{k+h}^2] + \frac{1}{2} \mathbb{E}[\tilde{e}_k^2] - \mathbb{E}[\tilde{e}_{k+h}\tilde{e}_k]
$$

we should obtain $\frac{1}{2} + \frac{1}{2} - 0 = 1$, which is indeed the case as seen in figure 2.58.
Figure 2.55: The observation errors for the VGSA model with a generic particle filter ...
Figure 2.56: ... and an extended particle filter.
Figure 2.57: The VGSA residuals Histogram. The residuals are fairly normal.
Figure 2.58: The VGSA residuals Variogram. The variogram is close to 1 as expected.
The observation is \( z_k = \ln S_{k+1} \)

\[
z_k = z_{k-1} + (\mu_S + \omega)\Delta t + \theta x_k + \sigma \sqrt{x_k} B_k
\]

with \( \omega = \frac{1}{\nu} \ln(1 - \theta \nu - \sigma^2 \nu/2) \).

and the hidden-state is

\[
x_k = Y_k(\Delta t)
\]

We could take advantage of the fact that VG provides an integrated density of stock return [182].

Calling \( z = \ln(S_k/S_{k-1}) \) and \( h = t_k - t_{k-1} \) and posing

\[
\xi_h = z - \mu_S h - \frac{h}{\nu} \ln(1 - \theta \nu - \sigma^2 \nu/2)
\]

we have

\[
p(z|h) = \frac{2 \exp(\theta \xi_h/\sigma^2)}{\nu \sqrt{2\pi \sigma^2}} \left( \frac{\xi_h^2}{2\sigma^2/\nu + \theta^2} \right)^{\frac{\nu}{2} - \frac{1}{2}} K_{\frac{\nu}{2} - \frac{1}{2}} \left( \frac{1}{\sigma^2} \sqrt{\xi_h^2 (2\sigma^2/\nu + \theta^2)} \right)
\]

where \( K_{\alpha}(x) \) corresponds to the modified Bessel function of second kind.

As we can see the dependence on the Gamma distribution is “integrated out” in the above. For the VGG for a given arrival rate \( dt^* = dY_t \) we have a VG distribution and

\[
d\ln S_t = (\mu + \omega)dt + B(\gamma(dt^*, 1, \nu); \theta, \sigma)
\]

and the corresponding integrated density becomes

\[
p(z|h, h^*) = \exp(\theta \xi_h/\sigma^2) \left( \frac{\xi_h^2}{2\sigma^2/\nu + \theta^2} \right)^{\frac{\nu}{2} - \frac{1}{2}} K_{\frac{\nu}{2} - \frac{1}{2}} \left( \frac{1}{\sigma^2} \sqrt{\xi_h^2 (2\sigma^2/\nu + \theta^2)} \right)
\]

Hence the idea of using the arrival-rate as the state and use the following algorithm:

1. **Initialize** the state \( x_0^{(i)} \) and the weight \( w_0^{(i)} \) for \( i \) between 1 and \( N_{\text{sims}} \)

2. **Simulate** the state \( x_k^{(i)} \) for \( i \) between 1 and \( N_{\text{sims}} \)

\[
\tilde{x}_k^{(i)} = F^{-1}(\mu_{a_i}, \nu_{a_i}; \Delta t, U^{(i)}[0, 1])
\]
where as before $F$ represents the Gamma CDF.

(3) Calculate the associated weights for each $i$

$$w_k^{(i)} = w_{k-1}^{(i)} p(z_k | \tilde{x}_k^{(i)})$$

with $p(z_k | \tilde{x}_k^{(i)})$ as defined in (2.32) where $h$ will be set to $\Delta t$ and $h^*$ to $\tilde{x}_k^{(i)}$

(4) Normalize the weights

$$\tilde{w}_k^{(i)} = \frac{w_k^{(i)}}{\sum_{i=1}^{N_{sims}} w_k^{(i)}}$$

(5) Resample the points $\tilde{x}_k^{(i)}$ and get $x_k^{(i)}$ and reset $w_k^{(i)} = \tilde{w}_k^{(i)} = 1/N_{sims}$.

(6) Increment $k$, Go back to step (2) and Stop at the end of the While loop.

As for VGSA, numeric tests were carried out in the following way:

After choosing a time-step $\Delta t = 1/252$, $\mu_S = 0$ and a parameter-set

$$\Psi = (\mu_a = 10.0, \nu_a = 0.01, \nu = 0.05, \sigma = 0.2, \theta = 0.002)$$

an artificial time-series of $N = 500$ spot-prices was generated. The above filtering algorithm was then applied to this time-series and the resulting likelihood was maximized. The optimal parameter-set was

$$\hat{\Psi} = (9.17, 0.19, 0.012, 0.21, 0.0019)$$

It therefore seems that the parameters $\nu$ and $\nu_a$ are not recovered properly. Hence the question how sensitive the observable spot-prices are to these variables? Simulating two time-series with the two different parameter sets, we can see in Figure 2.59 that the results could be very close.

This once again brings up the issue of inference reliability. Not having enough data points, we can get parameter-sets that are quite different from the real ones and that could generate similar time-series. This is consistent with what we have seen for diffusion-based processes.
Figure 2.59: Simulation of VGG-based Log-Stock-Prices with two different Parameter-Sets $\Psi = (\mu_a = 10.0, \nu_a = 0.01, \nu = 0.05, \sigma = 0.2, \theta = 0.002)$ and $\Psi = (9.17, 0.19, 0.012, 0.21, 0.0019)$. The observed time-series remain close.
A Bayesian Approach for VGSA

A similar approach as in the paragraph on the Bayesian approach for Heston could be used here, since the latent state follows the same square-root SDE. The only thing that changes, is the likelihood function. Instead of having a conditionally Log-Normal observation we have a conditionally VG observation. What is more, we do know the density of the VG distribution under a closed-form as previously mentioned.

Indeed as previously mentioned, we have the state (the arrival-rate)

\[ dy_t = \kappa(\theta - y_t)dt + \sigma_y \sqrt{y_t}dW_t \]

and the observation

\[ d\ln S_t = (\mu_S + \omega)dt + B(\gamma(dt^*, 1, \nu); \theta, \sigma) \]

and the corresponding conditional likelihood becomes

\[
p(\ln S_k | y_k, \Psi) = \frac{2 \exp(\theta x_h/\sigma^2)}{\nu \sigma \sqrt{2\pi} \Gamma(h)\nu^{-1/2}} \left( \frac{x_h^2}{2\sigma^2/\nu + \theta^2} \right)^{h^* \nu - \frac{1}{2}} K_{\frac{\nu}{2}} \left( \frac{1}{\sigma^2} \sqrt{x_h^2 (2\sigma^2/\nu + \theta^2)} \right)
\]

with \( K_\alpha(x) \) the modified Bessel function and

\[ x_h = \ln(S_k/S_{k-1}) - \mu_S h - \frac{h}{\nu} \ln(1 - \theta \nu - \sigma^2 \nu/2) \]

\[ h = \Delta t \]

and

\[ h^* = y_k \Delta t \]

Finally integrating over time, we have

\[
p(\ln S | y, \Psi) = \prod_{k=1}^{N} p(\ln S_k | y_k, \Psi)
\]

Note that in the classical VGSA model there is no correlation between the system noise and the observation noise. This means that the likelihood function will not depend on the parameters \( \kappa, \theta, \sigma_y \) and therefore the MH update step becomes almost a Gibbs sampler (except for the adjustment factor \( \sum_{k=1}^{N} x_k^2 \)).
2.4 Recapitulation

We tested three categories of models: the Heston/GARCH category where a pure diffusion assumption was used, the Bates category where Poisson jumps were added to the stock SDE, and the VG category where a Gamma distribution was applied to the time dimension.

2.4.1 Model Identification

We saw from the table 2.3.10 that in the pure diffusion category, a power of $3/2$ outperformed the Heston model (power of $1/2$). As stated, this is in line with the findings of Engle & Ishida [95].

Needless to say, adding Poisson jumps (Bates model) will reduce the MPE/RMSE of the filters, however will cause the number of parameters to increase. A simple comparison between the residual errors is therefore not fair.

In other words, given the fundamental differences between the categories, we need to judge their appropriateness not by comparing the residuals, but by using financial arguments such as Should the stock process contain jumps or not? Once a category is chosen, then we can compare the performance of models belonging to a given category.

Note that there exist a number of likelihood-based tools such as the Akaike Information Criterion [100], which will take into the account the number of parameters when assessing the goodness of fit for a model. These tools would therefore allow us to compare models belonging to different categories (e.g. Heston vs. VGSA). However these criteria remain valid only asymptotically. As we saw, this often requires a large number of data points, which may or may not have readily available.

2.4.2 Convergence Issues and Solutions

No matter which category we choose, it seems that the same convergence issues exist. For all the above models we can see that a parameter affecting the drift of the observation, is much easier to estimate than one affecting the noise of the observation.

For the pure diffusion category, we saw that all four parameters $\omega$, $\theta$, $\xi$ and $\rho$ were difficult to estimate (in some cases) and the two latter parameters, which affect the noise of the noise, were even harder to estimate properly.

For the Bates model, we saw that the jump parameters $\lambda$, $j$ were much more straightforward to estimate than the above mentioned four diffusion parameters.

For the VGSA models we saw that the VG parameters $\theta$, $\nu$ and $\sigma$ (which once again, affect the observation drift) are much easier to infer than the arrival-rate parameters $\kappa$, $\eta$ and $\lambda$. 

All this was explained via the *poor observability* at a daily frequency level, due to the fact that $\Delta t = o\left(\sqrt{\Delta t}\right)$. We tested the validity of this statement by artificially reducing the observation-noise and see the convergence rate increase dramatically.

As stated, a possible solution would be to employ more observation-points via the use of high-frequency data. We saw that the increase in the number of observations and the decrease in $\Delta t$ (after a certain level) do not cancel, and a higher frequency would indeed cause the likelihood function to have a higher value and provide a better estimation of the parameters.

In any case, since we do not know in advance how good the inference results are, and whether we are in the asymptotic area or not, it is always a good idea to perform a simulation test and determine the sampling distribution of each parameter.

In the next chapter, we shall apply these inference tools to a specific question: are the implied distributions from the stock and options markets consistent?
Chapter 3

The Consistency Problem

Whether cross-sectional option prices are consistent with the time-series properties of the underlying asset returns is probably the most fundamental of tests.

- David S. Bates

3.1 Introduction

In the previous chapter we discussed two different approaches for stochastic volatility parameter estimation: the cross-sectional one where we use a number of options prices for given strike prices (and possibly maturities); and the time-series approach where we use the stock prices over a certain period of time.

One natural question would therefore be the following: Will the theoretically invariant portion of the parameter-sets obtained by the two methods be the same?

More accurately, supposing we are at time $t = 0$ and we use $J$ options with strikes $K_1, ..., K_J$ and with maturity $T$, we have

$$
\hat{\Psi}_{\text{options}} = \arg\min_{\Psi} \sum_{j=1}^{J} [C_{\text{model}}(t = 0, S_0, K_j, T, \Psi) - C_{\text{mkt}}(t = 0, S_0, K_j, T)]^2
$$

(3.1)

the above options could include calls or puts.

On the other hand, during the period $[0, T]$ we can observe $(S_k)_{0 \leq k \leq N}$ corresponding to the time-points $t_0, ..., t_N$ with $t_0 = 0$ and $t_N = T$, and then apply one of the previously discussed Filters and estimate the parameter-set via the

---

1 Aït-Sahalia [6], Bakshi et al [20], Dumas et al. [88] have already asked a similar question, however they use a different approach for the Time-Series treatment.
Maximum-Likelihood method.

\[ \hat{\Psi}_{stocks} = \text{argmax}\{L(S_0, \ldots, S_N, \Psi)\} \] (3.2)

Now the question is how different these estimations for \((\xi, \rho)\) are and why?

As we saw in the previous chapter, the size of the time-interval \(\Delta t\) and the time-series length are to be questioned: Indeed \(\Delta t\) has to be small enough for us to be able to apply the Girsanov theorem. However we saw that for a very small \(\Delta t\), the filtering errors are so little that the MLE will not necessarily converge to the right parameter-set. On the other hand, we would need the time-series to be as long as possible, which requires a high observation frequency.

This brings up a more fundamental question: Current Financial Econometrics literature seems to make Inference-based conclusions using a limited amount of daily data. As we saw in the previous chapter, the time-series inference results are not necessarily reliable unless the number of observations is sufficiently large. This is the central question of this chapter: Are the implied parameters from the options markets and the assets time-series indeed inconsistent?

Many practical issues need to be questioned: how many strikes should we be using in the cross-sectional analysis and which ones? Should we use only OTM puts and calls for liquidity reasons? Many use a penalty function \(p()\) in the cross-sectional optimization in order to get reasonable results. Do we need such a function here?

In the cross-sectional method, what value for \(\nu_0\) are we using? Should we estimate this value together with the other four parameters? If so, should this estimated \(\hat{\nu}_0\) be used in the time-series?

If the results are substantially different for the parameters \(\xi\) and \(\rho\) (assuming the validity of the Girsanov theorem) can this test be used as an argument against the validity of the Heston stochastic volatility model? or would it mean that the options markets do not predict the stock movements as they should? And if so, does this mean that there is a profitable trading strategy to take? i.e. are options systematically mispriced?

If the Heston model is judged to be incorrect, what is the correct model? GARCH or \(3/2\)? Is the diffusion assumption itself to be questioned? Do we need to introduce jumps?

\(^2\)Note that an alternative method not involving any optimization would be a method of Matching of Moments. Indeed the Heston parameters \(\omega, \theta, \xi, \rho\) are analytically related to the first four moments of the time-series (mean, variance, skew, kurtosis). The calculation of the moments from the Time-series is fairly easy. The calculation of the moments from the options would require the use of the Carr-Madan [50] replication strategy using all available strike prices. However the information contained in the first four moments being less complete than the information contained in the density, the optimization method is more accurate.
Another way to approach the question is to reason in the following manner: If the information contained in the options markets is indeed inconsistent with the one embedded in the assets time-series, there should be a regularly and conclusively profitable trade strategy. For instance a higher volatility-of-volatility and more negative correlation in the options market should indicate the possibility of a profitable skewness trade (to be explained further below) in absence of crashes. We could therefore use the Profit/Loss of this trade as an empirical measure of the inconsistency of the information. If (and only if!) there exists a regular and definite profit generated from this strategy, we can conclude that there is inconsistency. It is important to note that this empirical measure is model free.

In our empirical analysis, unless stated otherwise, we shall use S&P500 Calls and Puts. There are two main reasons for this:

1. First of all these are the most liquid European Options available on CBOE. They expire on the third Friday of each contract month at the Open.

2. Also, there has already been abundant research carried out on these options. Indeed Aït-Sahalia [6], Bakshi, Cao and Chen [20], Bates [30], Dumas, Fleming and Whaley [88] and many others have all carried out their empirical analysis on S&P500 Options.

The data quality is obviously dependent on the degree of liquidity. Another issue we need to take into account is that of synchronization between the spot close-price and the option close-price. Indeed the timing of these two closings being off by a few minutes, the accuracy of the implied-volatility can be affected. Bates [32] specifically mentions this issue.

Let us be clear on the fact that this chapter does not constitute a thorough empirical study of the stock versus the options markets. It rather presents a set of examples of application of our inference tools constructed in the previous chapter. There clearly could be many other applications for these tools. As discussed in the second chapter, Model Identification is another instance.

It might seem that by avoiding the numeric optimization involved in our method we would gain precision, however given that the equations linking the first four moments and the four parameters are nonlinear, we would need to solve them numerically, which would be similar to an optimization.
3.2 The Consistency Test

In this section we shall compare the values of \((\xi, \rho)\) in the results \(\hat{\Psi}_{\text{options}}\) to \(\hat{\Psi}_{\text{stocks}}\) obtained via MLE. The time period \([0, T]\) is fixed and the time interval \(\Delta t\) for the stock is daily as in the previous chapter.

3.2.1 The Setting

The test is based upon SPX options as of 20020201 expiring in approximately 1 year from the calibration date. The daily time-series is taken during a period of 12 years corresponding to approximately 3000 points. The start of the period is 10 years before the calibration date and the end of the period is 1 year after the expiration of the options. Ideally we should only use the asset prices between the calibration date and the expiration to see whether the options predict the asset movements consistently. However this would provide us with too few observation points.

In what follows we will be considering one example of comparison between cross-sectional and time-series implied parameters. Many other similar examples were examined. They are not reported here since they do not change the conclusions.

The cross-sectional results (obtained from the options markets) belong to RBC Capital Markets and are similar to results reported by many others such as Bates [26] or Bakshi et al. [20]. The original contribution of our approach is presenting a systematic way to evaluate time-series embedded parameters. We shall do this via the methodologies\(^3\) detailed in the second chapter.

3.2.2 The Cross-Sectional Results

We consider one-year options as of January 2nd 2002 for close-to-the-money options. The calibration is done via LSE Monte-Carlo Mixing as well as the Fourier Inversion applied to the Heston model. We fix the instantaneous variance \(\nu_0\) at 0.04 and we take the index level at \(S_0 = 1154.67\) USD.

As usual we take the appropriate interest-rate \(r_T\) and dividend-yield\(^4\) \(q_T\) where \(T\) represents the options maturity. The dividend-yield could for instance be the one implied from the Forward contracts \(F_T\) calculated as

\[
q_T = r_T - 1/T \ln(F_T/S_0)
\]

\(^3\)These filter-based routines belong to the author and are available upon request.
\(^4\)No discrete dividends were considered.
We use various strike-price sets \((K_j)\) and determine the average optimal one-year parameters. Needless to say the results are obtained under the risk-neutral measure.

We obtain the risk-neutral implied parameter-set

\[
\begin{array}{cccc}
\omega & \theta & \xi & \rho \\
0.03620 & 1.1612 & 0.4202 & -0.6735 \\
\end{array}
\]

Table 3.1: Average Optimal Heston parameter-set (under the Risk-Neutral distribution) obtained via LSE applied to one-year SPX options on January 2002. Various Strike-Price sets were used.

which represents a rather high negative skewness and a high kurtosis.\(^5\) The long-term volatility is \(\sqrt{\omega/\theta} \approx 0.17\). Needless to say, these parameter values vary everyday, but usually remain in the same range.

**Robustness Issues for the Cross-Sectional Method**

1. For the Cross-Sectional analysis we have used a Mixing Monte-Carlo method. The Monte-Carlo time-steps of this method were spaced *weekly*. Therefore one natural question is how sensitive to this choice the results are. In order to verify this, we reran the simulations with *daily* Monte-Carlo time-steps and obtained

\[
\hat{\Psi}_{\text{options daily}} = (\omega = 0.036846, \theta = 1.169709, \xi = 0.42112, \rho = -0.67458)
\]

which is close to the original set.

We also checked the results with the volatility-of-volatility series method, as well as the Fourier inversion method, and obtained comparable parameters.

2. For our Cross-Sectional calibration we used Call Bid Prices. It is well known that Calls and Put prices are *not* always consistent. Indeed, as can be seen on figure 3.1 the Put and Call implied-volatilities are slightly different which seems to be a violation of Put-Call-Parity.\(^6\) However this difference is not large enough (The Put and Call Bid-Ask spreads actually overlap) and a profitable arbitrage cannot take place simply based upon this difference. This is why we consider the mid-point between Puts and Calls Bids and Asks.

\(^5\)We drop the “hat” notations for optimal parameters in this chapter for simplification. For example, instead of \(\hat{\omega}\) we simply write \(\omega\).

\(^6\)This is most probably due to the illiquidity of ITM options, as explained in [192].
Our implied-volatility is therefore

\[ \sigma_{imp} = \frac{1}{4} [\sigma_{imp}(Call_{Bid}) + \sigma_{imp}(Call_{Ask}) + \sigma_{imp}(Put_{Bid}) + \sigma_{imp}(Put_{Ask})] \]

Using these “mid” implied-volatilities as opposed to the original Call-Bids we obtain a parameter-set

\[ \Psi_{options-\text{mid-call-put}} = (\omega = 0.043184, \theta = 1.173119, \xi = 0.40258, \rho = -0.64593) \]
3. If we do include \( v_0 \) in the set of parameters \( \Psi = (\omega, \theta, \xi, \rho, v_0) \), then we obtain

\[
\hat{\Psi}_{\text{options - mid-call-put}} = (\omega = 0.043224, \theta = 1.144957, \xi = 0.482009, \rho = -0.661427, \sqrt{v_0} = 0.224659)
\]

It is possible to see that the optimal \( \sqrt{v_0} \) is around 0.20 which corresponds to our initial choice.

4. As already mentioned, further from the money options are less reliable in terms of pricing and liquidity. However disregarding them decreases the cross-sectional sensitivity to the volatility-of-volatility parameter. Adding to the previous close-to-the-money strikes, additional further-from-the-money ones, we find

\[
\hat{\Psi}_{\text{options}} = (\omega = 0.035896, \theta = 1.149324, \xi = 0.386453, \rho = -0.659319, \sqrt{v_0} = 0.221988)
\]

Again, the drift parameters are stable, and so is \( v_0 \). The question is how the volatility parameters are affected? Interestingly we do not observe a great difference from what we had with the previous sets. We therefore have a good degree of robustness.

In any case, we use various sets of strike-prices and take an average over the optimal parameter-sets.

5. One issue to consider in the Cross-Sectional method is how the risk-neutral implied distribution, or in our case the parameter-set \( \Psi \) evolves in time. Needless to say, if the model was perfectly correct these parameters would never change, however as we know this is never the case. The question therefore becomes how time-homogeneous these parameters are.

Considering the same maturity 20021221 but at a date closer to this maturity, we use close-to-the-money strikes. More accurately we stand at 20020903, take the spot at 878.02 USD, and use the yield-curve as of 20020903.

The strikes are

\[
K_{\text{set}} = \{775.00, 800.00, 825.00, 850.00, 875.00, 900.00, 925.00, 950.00, 975.00\}
\]

The optimization via Monte-Carlo mixing provides:

\[
\hat{\Psi}_{\text{options}} = (\omega = 0.0501244, \theta = 1.189817, \xi = 0.547149, \rho = -0.661552, \sqrt{v_0} = 0.265441)
\]

which is not too far from the other parameter-sets.
3.2.3 Time-Series Results

As mentioned, the first idea is to choose a period corresponding to the life of the options considered in the previous section. Indeed we would like to see whether the options are predicting the underlying asset dynamics correctly during their life. However this provides us with one year of daily data or 252 points which as we know from the previous chapter is highly insufficient for time-series estimators.

In order to obtain more reliable results we use various Filters (EKF, EPF ...) and take the average optimal parameter-set. For a period of 12 years ending on January 2004 (which includes the options life) applying the filters studied in the previous chapter, we obtain the following average results:

<table>
<thead>
<tr>
<th>ω</th>
<th>θ</th>
<th>ξ</th>
<th>ρ</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.018620</td>
<td>0.523947</td>
<td>0.096389</td>
<td>-0.132527</td>
</tr>
</tbody>
</table>

Table 3.2: Average Optimal Heston parameter-set (under the Statistical distribution) obtained via Filtered MLE applied to SPX between January 1992 and January 2004. Various Filters were used in the MLE.

The above results show a lower (ξ, ρ) and therefore a lower implied skewness and kurtosis, than the ones obtained from the options markets.

Robustness Issues for the Time-Series Method

Given the above results, it would be instructive to test the sensitivity of the observations to the drift parameters (ω, θ) on the one hand, and to the volatility parameters (ξ, ρ) on the other.\(^7\)

The point is that even if the state \(v_k\) itself is greatly affected by these volatility parameters, the impact of these parameters on the observations is small. However the impact of the drift parameters is quite large. This could explain why the Cross-Sectional and Time-Series volatility-of-volatility parameters are not close.

This point can be observed in the simulations represented in figures 3.2 to 3.5. Note that this issue is related to the discussion in Chapter 2 on the Sampling Distribution. Indeed as previously stated, ξ and ρ have a lesser effect on the observations since they affect the “noise of the noise”.

\(^7\)Note that we could not have done this separation in a nonparametric model such as [6].
The observations have little sensitivity to the volatility parameters. One year simulation with $\sqrt{v_0} = 0.20$, $\omega = 0.04$, $\theta = 0.5$. Cross-Sectional uses $\xi = 0.036$ and $\rho = 0.50$ while Time-Series uses $\xi = 0.09$ and $\rho = -0.80$. This is consistent with what we had seen previously.
Figure 3.3: The state has a great deal of sensitivity to the volatility parameters. One year simulation with $\sqrt{v_0} = 0.20$, $\omega = 0.04$, $\theta = 0.5$. Cross-Sectional uses $\xi = 0.036$ and $\rho = 0.50$ while Time-Series uses $\xi = 0.09$ and $\rho = -0.80$. 
Figure 3.4: The observations have a great deal of sensitivity to the drift parameters. One year simulation with $\sqrt{v_0} = 0.20$, $\xi = 0.036$, $\rho = 0.50$. Cross-Sectional uses $\omega = 0.04$ and $\theta = 0.50$ while Time-Series uses $\omega = 0.08$ and $\theta = 5.0$. 
Figure 3.5: The state has a great deal of sensitivity to the drift parameters. One year simulation with $\sqrt{v_0} = 0.20$, $\xi = 0.036$, $\rho = 0.50$. Cross-Sectional uses $\omega = 0.04$ and $\theta = 0.50$ while Time-Series uses $\omega = 0.08$ and $\theta = 5.0$. 
3.2.4 Financial Interpretation

The current Financial Econometrics consensus is the following: No matter which case we consider, the cross-sectional parameters $\xi$ and $\rho$ are always greater (in absolute-value) than the time-series ones.

This means that the skewness and the kurtosis implied from options are stronger than those implied from the time-series. As we will see in the following paragraphs, this could suggest a trade to take advantage of this inconsistency, supposing that the options are misjudging the spot movements.

We can observe the above statement graphically by plotting the SPX volatility smile from the options market prices on the one hand, and from the time-series implied parameters on the other. Note that we need no calibration for the options since we are using the usual Black-Scholes implied volatility.

Figure 3.6 shows the difference between the two slopes. Again the options curve has a stronger (negative) slope, which is consistent with a stronger negative product $\xi\rho$.

As explained in [69] the higher moments of the stock-price return can be calculated from the stochastic-volatility model parameters.

Indeed, for a given parameter-set $\Psi = (\omega, \theta, \xi, \rho)$ we have

$$\text{skewness} = \left( \frac{3\xi\rho e^{\frac{\rho}{2}\theta t}}{\sqrt{\theta}} \right) \left[ \frac{\theta}{\pi(1 - \theta T + \theta Te^{\theta T})} \left( \frac{\pi}{(1 - \theta T + \theta Te^{\theta T}) + v_0(e^{\theta T} - 1)} \right) \right]$$

and

$$\text{kurtosis} = 3 \left[ 1 + \xi^2 \left( \frac{\pi A_1 - v_0 A_2}{B} \right) \right]$$

with $y = \theta T$ and

$$A_1 = [1 + 4e^y - 5e^{2y} + 4ye^{2y} + 2ye^{2y}] + 4\rho^2[6e^y - 6e^{2y} + 4ye^y + 2ye^y + y^2e^y]$$

$$A_2 = 2[1 - e^{2y} + 2ye^y] + 8\rho^2[2e^y - 2e^{2y} + 2ye^y + y^2e^y]$$

$$B = 2\theta \left( \frac{\omega}{\theta} - ye^y + v_0(e^y - 1) \right)^2$$

Without entering into the details of the calculations, we can see that for given $\omega$ and $\theta$, higher $(\xi, |\rho|)$ correspond to higher skewness and kurtosis. As we said in the previous chapter, the skewness depends on $\omega$, $\theta$ and the product $\xi\rho$ which has a more reliable estimation than the separate values of $\xi$ and $\rho$. This makes the estimation of the skewness more trustworthy.
Figure 3.6: Comparing SPX Cross-Sectional and Time-Series Volatility Smiles (with historic $\xi$ and $\rho$) as of Jan 2nd 2002. The spot is at 1154.67 USD.
3.3 The “Peso” Theory

3.3.1 Background

As [6] mentions, one possibility regarding the Cross-Sectional vs. Time-Series observed differences is the following:

As we know the Time-Series corresponds to one realization of the stock-return stochastic process. Now supposing that the true stock Stochastic Differential Equation (SDE) contains jumps, there is a possibility that the historic path we are observing does not contain any of these jumps. This is referred to as the Peso theory.

As mentioned in [12], this term goes back to Milton Friedman in his analysis of Mexican Peso during the early 70’s. The Mexican interest rates remained significantly above the US interest rates, although the peso was pegged at 0.08 dollars per peso. Friedman argued that the interest rates reflected an expectation about a future devaluation of the peso. In August 1976, the peso was devaluated by 37.5% to a new rate of 0.05 dollar per peso, thus validating the previous interest rates differential.

This assumption seems reasonable since as we saw in the previous tests, the Cross-Sectional method usually provides higher volatility parameters ($\xi, \rho$) and therefore higher skewness (in absolute value) and kurtosis. Introducing a jump component in the options pricing model should lower these optimal parameters.

Note that in [3], Aït-Sahalia tries to find out whether the discrete observations of S&P500 come from a diffusion, or from a distribution containing jumps? He derives a criteria for continuity of the paths

$$\frac{\partial^2}{\partial x \partial y} \ln (p(\Delta t, y = X_{t+\Delta t} | x = X_t)) > 0$$

for every $\Delta t > 0$ and given $(x, y)$.

Based upon the implied cross-sectional distribution he finds that S&P500 options do consider jumps in the paths.

Using the Jump Diffusion model as we did in the previous chapter

$$d \ln S_t = (\mu_S - \frac{1}{2} \nu_t + \lambda_j) dt + \sqrt{\nu_t} dB_t + \ln(1 - j) dN_t$$

$$d \nu_t = (\omega - \theta \nu_t) dt + \xi \sqrt{\nu_t} dZ_t$$

we may very well see no difference introduced from the parameters ($\lambda, j$) for the Time-Series and we can even disregard them. However, this does not mean that the stock process does not contain jumps, but rather this specific path happens

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Jackwerth and Rubinstein [155] refer to this phenomena as Crash-o-Phobia.
to contain none. The Options on the other hand, always include the possibility of jumps in their pricing. Adding \((\lambda, j)\) will affect the resulting \((\xi, \rho)\) from the Cross-Sectional method.

3.3.2 Numeric Results

We use the same Options and Time-Series as in the previous section. As shown in Merton’s paper [190] we have for a given volatility path \(\sigma = \sqrt{v}\)

\[
Call = \sum_{n=0}^{\infty} e^{-\lambda(1-j)T} \frac{(\lambda(1-j)T)^n}{n!} C_{BS}(S, K, T, \sigma, r_n)
\]

with

\[
r_n = r + \lambda j + \frac{n}{T} \ln(1 - j)
\]

We then take the expectation upon the volatility stochastic process as we usually do in a Mixing algorithm.\(^9\)

We find for the parameter-set \(\hat{\Psi} = (\omega, \theta, \xi, \rho, \lambda, j)\) the values

\[
\hat{\Psi}_{\text{options}} = (\omega, \theta, \xi, \rho, \sqrt{v_0}, \lambda, j) = (0.032648, 1.165598, 0.360646, -0.585302, 0.218333, 0.008982, 0.913772)
\]

instead of the previous pure-diffusion parameter-set

\[
\hat{\Psi}_{\text{options - mid - call - put}} = (\omega = 0.043224, \theta = 1.144957, \xi = 0.482009, \rho = -0.661427, \sqrt{v_0} = 0.224659)
\]

As we see even with the addition of jump parameters \((\lambda, j)\), the cross-sectional volatility parameters \((\xi, \rho)\) remain significantly above the time-series ones. This is in agreement with the findings of Bakshi, Cao and Chen [20]. We have a small \(\lambda\) and a \(j\) close to one. This means that options are expecting a large but infrequent jump, \textit{i.e.} they are factoring in the possibility of a crash.

\[^9\text{Note that an alternative method would be to use a Fourier inversion of the known Characteristic Function as Lewis does in [178] or [180].}\]
3.4 Trading Strategies

Supposing that the model we are dealing with is correct, if the options are mistaken in evaluating the stock distribution during their lifetime, there should be an arbitrage opportunity to take advantage of.

The ninth chapter of the Härdle et al. book [128] has a description of these strategies. Note that both these strategies are European and cannot be changed\textsuperscript{10} until maturity.

At this point we should reiterate that the Profit and Loss of this trade could be used as an empirical and model-free measure of how consistent or inconsistent the information embedded in the options is with the one in the underlying stocks.

3.4.1 Skewness Trades

To capture an undervalued third moment we can buy OTM calls and sell OTM puts. Note that Aït-Sahalia [6] says that the options are overly skewed, which means that the options skew is larger in absolute value. However given the negative sign of the skew, the cross-sectional skew is actually lower than the one implied by the Time-Series, hence the described strategy.

Note that in order to be immune to parallel-shifts of the volatility-curve, we should make the trade as vega-neutral as possible. The correspondence between the Call and the Put is usually not one-to-one.

Therefore calling $V$ the vega, $\Delta$'s the hedge-ratios for $C$ the call and $P$ the put option, the hedged portfolio $\Pi$ will be

$$\Pi = C(S_t, K_C) - \frac{V_C}{V_P} P(S_t, K_P) - \left( \Delta_C - \frac{V_C}{V_P} \Delta_P \right) S_t$$

and the positions in the options should be dynamically adjusted in theory. However that would cause too much transaction cost and exposure to the bid-ask spread.

As we shall see in the paragraph on “Exact Replication” there exist more elaborate strategies, better exploiting the third moment differences.

3.4.2 Kurtosis Trades

To capture an overvalued fourth moment, we need to use the “fat tails” of the distribution. For this we can for instance sell ATM and far OTM options, and buy close OTM options.

\textsuperscript{10} As we will see further, we could unwind the deal prior to expiration. However we would then be subject to the movements of options prices.
3.4.3 Directional Risks

Despite the Delta-Hedging, the skewness trade applied to an undervalued third moment has an exposure to the direction of the markets. A bullish market is favorable to it, and a bearish one unfavorable.

The Kurtosis trade applied to an overvalued fourth moment generates a profit if the market stays at one point and moves sideways but loses money if there are large movements.

This exposure to market conditions is consistent with the Peso theory. The Skewness and Kurtosis trading strategies above are profitable given the options implied moments, unless the options were actually right in factoring in a large and sudden downward movement. This also makes sense since the way the options were priced changed only after the crash of 1987. Prior to that, the volatility negative-skew was practically absent altogether. Figures 3.7 and 3.8 show generic examples of the strategies described above.

Note that as the skew formula in [69] shows, the volatility of volatility $\xi$ affects the skew as much as the correlation $\rho$ does. This explains why sudden upward movements can hurt us as well. If the overall correlation is negative but there are large movements in both directions we will have large third (in absolute value) and fourth moments, which would make the options expectations correct. Indeed as we will see in the following example, a large upward movement can make us lose on our hedge account.

As many such as [32], [128] have mentioned, it is possible to interpret this trade as an Insurance Selling strategy. The trade will generate moderate and consistent profits if no crash happens. But if the crash does happen we could suffer a large loss.

Skewness vs. Kurtosis

The Skewness Trade seems to be a simpler one and has a better chance to be realized. Indeed in order to have a large negative skew, we need a large volatility-of-volatility $\xi$ (as we do for the kurtosis trade) and a large negative correlation $\rho$.

In other words, if for a given stock time-series we have a large volatility-of-volatility but a weak correlation, we will not have a kurtosis trade opportunity but we will have a Skewness Trade opportunity. Indeed the historic skew will be small and the historic kurtosis high.

Graphically we could have the following interpretation: For these assets, the Historic distribution does have fat tails, but remains symmetric. While the Implied distribution has a fatter left tail. This is why we have a Skewness Trade opportunity, even if we do not have a Kurtosis trade opportunity.

Finally, as we previously mentioned the estimation of the skewness from a time-
The Volatility Process

series is more reliable since it only depends on the \textit{product} of the volatility-of-volatility and the correlation.

### 3.4.4 An Exact Replication

These trading strategies can be refined using a Carr-Madan replication. Indeed as explained in [50] we have for any payoff function \( f() \) the following identity

\[
E[f(S_T)] = f(F) + e^{rT} \int_0^F f''(K)P(S_0, K, t = 0, T)dK + e^{rT} \int_F^{+\infty} f''(K)C(S_0, K, t = 0, T)dK
\]

with \( F = S_0e^{rT} \) the forward price.

In order to get the Das [69] skew and kurtosis calculations, we need to take for the \( n^{th} \) moment

\[ f(S_T) = (Z_T - E(Z_T))^n \]

with

\[ Z_T = \ln(S_T/S_0) \]

However this trade will clearly have a much higher transaction cost than the one described in the previous paragraph.
Figure 3.7: A generic example of a Skewness strategy to take advantage of the undervaluation of the skew by Options. This strategy could be improved upon by trading additional OTM puts and calls.
Figure 3.8: A generic example of a Kurtosis strategy to take advantage of the overvaluation of the kurtosis by Options. This strategy could be improved upon by trading additional puts and calls.
3.4.5 The Mirror Trades

Should we see the opposite conditions in the market, \textit{i.e.} having the skew (in absolute value) or kurtosis \textit{undervalued} by the options given a historic path, we could obviously put on the mirror trades.

The inverse of the Peso theory would be as follows: the stock in question has already had a crash and the options are supposing there probably will \textit{not} be another one in the near future.

Setting up the overvalued kurtosis trade in the previous paragraph, we picked up a premium and made an immediate profit and hoped that there would not be a sudden movement.

Here we start by losing money and hope a crash will happen within the life of the option so we can generate a profit. Jumps and crashes being rare by nature, this trade does not seem very attractive.

What is more if there was a recent crash, the possibility of another one is indeed reduced and we should believe the options prediction. However these mirror trades could be considered as \textit{buying insurance} and therefore as a protection against a possible crash.
3.4.6 An Example of the Skewness Trade

The algorithm is as follows:
For a given date \( t_0 \) we have \( S_0 \) and choose the closest maturity to \( T = t_0 + 0.25 \) in order to have a three month trade, we then take the Call and Put Strikes \( K_C \) and \( K_P \) as the closest ones to 1.10\( S_0 \) and 0.90\( S_0 \) respectively.

The original cost is therefore

\[
\text{options}(0) = \text{Call}_{\text{Ask}}(S_0, K_C, t_0, T) - \text{Put}_{\text{Bid}}(S_0, K_P, t_0, T)
\]

Indeed we buy a Call at the Offer price and sell the Put at the market Bid price. At maturity, the position is worth

\[
\text{options}(T) = \text{MAX}(0, S_T - K_C) - \text{MAX}(0, K_P - S_T)
\]

During the trade we have a Delta hedging cash-flow of

\[
hedge = -\sum_{t=0}^{T-1} \Delta(S_t, t, T)(S_{t+1} - S_t)
\]

with

\[
\Delta(S_t, t, T) = \Delta_{\text{Call}}(S_t, K_C, t, T, \sigma_{\text{imp}}(t_0, K_C)) - \Delta_{\text{Put}}(S_t, K_P, t, T, \sigma_{\text{imp}}(t_0, K_P))
\]

where the above implied-volatilities used in the hedge-ratios are using the Mid prices (between Bid and Ask prices),

as well as an interest-rate cash-flow of

\[
\text{interest} = \sum_{t=0}^{T-1} \Delta(S_t, t, T)(e^{r\Delta t} - 1)S_t
\]

Our final profit or loss (PnL) is therefore

\[
PnL = \text{options}(T) - \text{options}(0) + hedge + interest
\]

If the options implied skew is indeed higher than justified by the stock movements, then this trade should be profitable. However in case of a sudden large movement, this will not be true anymore.
Figure 3.9: Historic spot level movements during the trade period.
Figure 3.10: Hedging PnL generated during the trade period. As we can see, losses occur upon jumps.
Figure 3.11: Cumulative hedging PnL generated during the trade period. This positive PnL will be offset by the option premiums and payoffs.
We consider the case of S&P500 between 20020404 and 20020622. At that point in time $S_0 = 1126.34$ USD, which means we can take $K_C = 1250$ USD and $K_P = 1050$ USD.

We also have $Call_{Ask}(t_0, K = 1250) = 3.20$ USD and $Put_{Bid}(t_0, K = 1050) = 14.20$ USD, as well as the mid implied volatilities of $\sigma_{imp}(K_C) = 0.154$ and $\sigma_{imp}(K_P) = 0.214$.

As can be seen in figures 3.9 and 3.10, the sudden spot movements generate most of the loss (for instance around day 50). We have at the end of the trade in USD’s

\[
S_T = 989.14 \\
hedge = 50.39 \\
interest = 1.32
\]

Therefore the final PnL in USD is\(^{11}\)

\[
PnL = [0 - (1050 - 989.14)] + (14.20 - 3.20) + 50.39 + 1.32 = 1.85
\]

As we can see, we hardly generated a profit, given the “jumps” occurring in the middle of the trade. Note that we generated a profit in the beginning by selling an OTM Put that was more expensive than the OTM Call we bought. We lost a large amount because the spot ended up below the Put strike. However we compensated that via the hedge.

The Options Bid-Ask Spread

It is important to know where we are buying the Call and Selling the Put on the start-date. Are we buying the Call at the Offer-price and selling the Put at the Bid-Price? In which case we can lose the Bid-Ask spread, comparing to the case where we would buy and sell both options at the Mid-Market. This spread is in average around 1 USD for ten-percent OTM SPX options.

Early Termination

We also should consider an early unwinding of the trade: Indeed as we get closer to maturity, our hedge-ratio might be close to one, which will make our hedge-account extremely sensitive to adverse stock movements. In order to have a smoother PnL, we can buy back the Put and sell the Call at a date (e.g. one

\(^{11}\)Proprietary results generated within RBC Capital Markets.
month) prior to maturity. Again, it is important to know whether we are unwinding the trade by selling the Call at the Bid and buying back the Put at the Offer? If so, we will have suffered from the Bid-Ask spread twice: once on the start-date and once on the unwinding (termination) date.

This is not just a small detail, indeed having the right execution (at mid-market) can change the average sign of the PnL altogether. What is more, regardless of the Bid-Ask spread, we are subject to the movements of the options prices. Whereas if we hold the positions until expiration, we will have a pure strategy between the original options prices and the spot price movements.

**Implied Volatility Term-Structure**

Yet another issue to take into account is that in our back-testing, we used fixed implied-volatilities in order to calculate the hedge-ratios during the life of the trade. In reality the implied-volatilities change every day even if we assume a Sticky-Strike regime where the stock-price will not affect the implied-volatility level. Indeed even if our Strikes are fixed, the time-to-maturities of the options decrease, and this will make the implied-volatilities vary. For S&P500 the term-structure of implied-volatility is upward-sloping which means that theoretically all implied-volatilities should come down from their original levels at the unwinding date.

**Which Hedge-Ratio should we use?**

In the hedging of our skew portfolio, which $\Delta$ should we apply? In other words, the question is which implied-volatility we should use in the usual

$$e^{-q(T-t)}N\left(d_1(S_t, K, t, T, \sigma_{imp})\right)$$

If we believe that the volatility predicted by the options is wrong and the historic levels are correct we should then use

$$\sigma_{stocks}(K, T) = C^{-1}_{BS}(C_{model}(S_0, t_0, K, T, \hat{\Psi}_{ stocks}))$$

where

$$\hat{\Psi}_{ stocks} = (\hat{\omega}_{ opts}, \hat{\theta}_{ opts}, \hat{\xi}_{ stocks}, \hat{\rho}_{ stocks})$$

Note that this might give us a mismatch in terms of mark-to-market with the existing option levels in the market. However if the Time-Series is really correct, the skew should eventually **collapse** before the options mature.\(^{12}\) We should note

\(^{12}\)Bates [29] suggests the use of an adjusted Delta as

$$\Delta \approx \Delta_{BS} - \frac{K}{S} \frac{\partial \sigma}{\partial K}$$
however that using the options implied volatilities makes better practical sense, since those are the ones where the options are actually traded.

### 3.4.7 Multiple Trades

The next natural step would be to repeat the previous trade in order to see whether the trade would be statistically profitable. We use SPX Puts and Calls between 20020102 and 20030201 on the expiration month such that the original life of the trade is around three months. We systematically unwind the trades around 20 business days to expiration. Once again, we buy ten-percent OTM Calls and we sell ten-percent OTM Puts. We cover in this manner forty different cases. We calculate the PnL’s as previously described\(^\text{13}\) and take their average.

*The results are mixed: If we put the trade on and unwind at the Bid and Ask levels we will actually suffer a loss. However if we can execute at the Mid, then we will generate a profit.*

This shows a lack of decisive proof on an inconsistency between the options and stock markets\(^\text{14}\). Indeed we have used the PnL of this trade as a measure of discrepancy\(^\text{15}\).

\(^{13}\)Proprietary results generated within RBC Capital Markets.

\(^{14}\)Note that this trade generates a regular and stable profit and sudden large losses. This is in agreement with the interpretation of *selling insurance* and collecting the premiums. It is very profitable until there is a “disaster”.

\(^{15}\)There is a case where a skew trade should be considered. Even if we have an inefficient estimate of ξ and ρ, we do have their Sampling Distributions, as seen in the previous chapter. If for instance the average estimate of ξ is 0.03, supposing the lowest and highest estimates are respectively −0.20 and 0.20, and if ξ\(_{opt}\) = 0.40 then there is an inconsistency in a conclusive manner. Indeed we would then have our cross-sectional volatility-of-volatility far superior to its highest possible time-series estimate.

where \( V \) represents the option vega.

However as he points out, this is the hedge-ratio as perceived by the options market and this perception could very well be wrong. After all this is what we are trying to take advantage of: the mispricing of the skew by the options, supposing that the historic time-series has the same dynamics as the future spot movements.

\(^{13}\)Proprietary results generated within RBC Capital Markets.

\(^{14}\)Note that this trade generates a regular and stable profit and sudden large losses. This is in agreement with the interpretation of *selling insurance* and collecting the premiums. It is very profitable until there is a “disaster”.

\(^{15}\)There is a case where a skew trade should be considered. Even if we have an inefficient estimate of ξ and ρ, we do have their Sampling Distributions, as seen in the previous chapter. If for instance the average estimate of ξ is 0.03, supposing the lowest and highest estimates are respectively −0.20 and 0.20, and if ξ\(_{opt}\) = 0.40 then there is an inconsistency in a conclusive manner. Indeed we would then have our cross-sectional volatility-of-volatility far superior to its highest possible time-series estimate.
3.4.8 High Volatility-of-Volatility and High Correlation

As previously discussed many stocks do have a high historic volatility-of-volatility $\xi$ however given a weak (or even positive) spot-volatility correlation $\rho$, the historic skew is still very low. This is specially true of “penny” stocks. Indeed when these stocks increase in price, in some sense they “come back to life” and therefore become more volatile. This means that the historic skew is actually positive, which seems to indicate an even stronger case for a Skewness Trade. However given that we are dealing with penny stocks the possibility of a crash for these stocks is high and that is precisely what causes the negative option-implied skew! The stock GW (Grey Wolf Inc.) in figure 3.14 is a good example for this case. This presents a trading opportunity as shown in figure 3.15. On the other hand, there are cases such as MSFT (Microsoft) where we do have a strong historic negative correlation as well as a high volatility-of-volatility. As the stock price goes down, the asset becomes riskier and therefore more volatile. We can see this in figure 3.16. This justifies the option-implied skew observable in figure 3.17 and means there is no trade opportunity. The safest trade therefore seems to be an Index Skewness Trade given that the likelihood of a crash is lower thanks to the diversification effect.

Note that the strong negative historic skew is not limited to individual stocks. Indeed taking the case of the NDX index in figures 3.18 and 3.19, we can see that there is no trading opportunity available and the historic skewness is in line with the one implied by the options prices.

There are therefore two different possible reasons why we may have a Skewness Trade Opportunity:

1. Weak Historic Volatility-of-Volatility (e.g. SPX (S&P500))
2. Weak Historic Correlation (e.g. GW (Grey Wolf Inc.))

If none of the above is verified (e.g. NDX (Nasdaq) or MSFT (Microsoft)) there is no skew trading opportunity.

The graphical interpretation seen in the following figures is based upon the comparison of the observable options-implied skew

$$\sigma_{\text{options}}^{imp}(K, T) = C_{BS}^{-1}(C_{mkt}(S_0, t_0, K, T))$$

and the skew implied from historic stock-price movements

$$\sigma_{\text{stocks}}^{imp}(K, T) = C_{BS}^{-1}\left(C_{model}(S_0, t_0, K, T, \hat{\Psi}_{\text{stocks}})\right)$$

where $C_{BS}$ corresponds the usual Black-Scholes pricing function.

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16These tests were performed around end of March 2003.
Again, we use the option-implied volatility-drift parameters $\hat{\sigma}_{\text{options}}, \hat{\theta}_{\text{options}}$ in $\Psi_{\text{stocks}}$. The only assumption here would be that of Diffusion in the processes. Indeed then according to the Girsanov theorem the volatility-of-volatility and the correlation should be the same for the continuous statistical and risk-neutral processes.
Figure 3.12: A Strong Option-Implied Skew: Comparing MMM (3M Co) Cross-Sectional and Time-Series Volatility Smiles as of Mar 28 2003. The Spot is at 131.66 USD.
Figure 3.13: A Weak Option-Implied Skew: Comparing CMI (Cummins Inc) Cross-Sectional and Time-Series Volatility Smiles as of Mar 28 2003. The Spot is at 24.59 USD.
Figure 3.14: GW (Grey Wolf Inc.) Historic prices (20020331-20030331) show a high volatility-of-volatility but a weak stock-volatility correlation. The resulting negative skew is low.
Figure 3.15: The Historic GW (Grey Wolf Inc.) Skew is low and not in agreement with the options prices. There is a skew trading opportunity here.
Figure 3.16: MSFT (Microsoft) Historic prices (20020331-20030331) show a high volatility-of-volatility and a strong negative stock-volatility correlation. The resulting negative skew is high.
Figure 3.17: The Historic MSFT (Microsoft) Skew is high and in agreement with the options prices. There is no skew trading opportunity here.
Figure 3.18: NDX (Nasdaq) Historic prices (20020331-20030331) show a high volatility-of-volatility and a strong negative stock-volatility correlation. The resulting negative skew is high.
Figure 3.19: The Historic NDX (Nasdaq) Skew is high and in agreement with the options prices. There is no skew trading opportunity here.
3.5 Non-Gaussian Case

As previously discussed, once we start dealing with some of the pure-jump models such as VGG, we will not have the Girsanov theorem anymore and cannot compare the parameters directly. However, no matter what the arrival process is, we still have the VG parameters $(\sigma, \nu, \theta)$ as in

$$d\ln S_t = (\mu_S + \omega)dt + X(dt; \sigma, \nu, \theta)$$

where as before $\mu_S$ is the real-world statistical drift of the stock log-return and $\omega = \frac{1}{\nu} \ln(1 - \theta \nu - \sigma^2 \nu/2)$.

As for $X(dt; \sigma, \nu, \theta)$ it has the following meaning:

$$X(dt; \sigma, \nu, \theta) = B(\gamma(dt, 1, \nu); \theta, \sigma)$$

where $B(dt; \theta, \sigma)$ would be a Brownian Motion with drift $\theta$ and volatility $\sigma$. In other words

$$B(dt; \theta, \sigma) = \theta dt + \sigma \sqrt{dt} N(0, 1)$$

where $N(0, 1)$ is a standard Gaussian realization.

What is more, we know what the centralized third and fourth moments (Skewness and Kurtosis) are

skewness = $(2\theta^3 \nu^2 + 3\sigma^2 \theta \nu)t$

kurtosis = $(3\sigma^4 \nu + 12\sigma^2 \theta^2 \nu^2 + 6\theta^4 \nu^3)t + (3\sigma^4 + 6\sigma^2 \theta^2 \nu + 3\theta^4 \nu^2)t^2$

We therefore can always compare the skewness and kurtosis implied from time-series with those implied from options. However a mismatch between the two does not indicate an arbitrage opportunity since once again we are comparing them under two different measures.

Having said this, the determination of the statistical density $p()$ and the risk-neutral density $q()$ is still useful in the sense that it could allow us to determine the optimal position we would take in the derivatives market given a Utility function, as described in [52], [53].

Indeed having an increasing concave utility function $U()$, the idea is to find the optimal payoff $\phi(S)$ maximizing the expected utility at a given horizon $T$, among all possible payoffs $f(S)$

$$\phi = \arg\max \int_0^{+\infty} U[f(S_T)] p(S_T) dS_T$$

In addition to this, we have the initial budget $W_0$ which imposes a constraint: the discounted risk-neutral expected value of the payoff cannot be greater than this initial budget:

$$\exp(-rT) \int_0^{+\infty} f(S_T) q(S_T) dS_T \leq W_0$$
This can be seen by using a “self-financing” portfolio argument as was done by Black and Scholes. Using the two equations above we can write the Lagrangian

\[ \mathcal{L}(f) = \int_0^{+\infty} U[f(S_T)]p(S_T)dS_T - \lambda \exp(-rT) \int_0^{+\infty} f(S_T)q(S_T)dS_T \]

where \( \lambda \) is the Lagrange multiplier. We then can differentiate with respect to the payoff \( f() \) and obtain the optimal payoff satisfying

\[ \exp(rT) \frac{p(S)}{q(S)} U'[\phi(S)] = \lambda \]

or equivalently

\[ \phi(S) = (U')^{-1} \left( \lambda \exp(-rT) \frac{q(S)}{p(S)} \right) \]

and the constant \( \lambda \) could be determined by a normalization such as

\[ \exp(-rT) \int_0^{+\infty} (U')^{-1} \left( \lambda \exp(-rT) \frac{q(S)}{p(S)} \right) q(S_T)dS_T = W_0 \]

This would provide us with the optimal payoff function that we would need to choose in the derivatives market, and therefore motivates the estimation of the statistical and risk-neutral densities \( p \) and \( q \) even for the non-Gaussian case.

### 3.5.1 VGSA

Unlike VGG and many other pure-jump models, VGSA has a conditionally Gaussian arrival rate. This means that the volatility of the arrival-rate \( \lambda \) should remain the same under the statistical and risk-neutral measures.

We therefore do have an analog approach to the diffusion based models for VGSA.

#### VGSA vs. VG

In their original paper [182] Carr, Madan and Chang find comparable results for the VG model applied to S&P500 for 1992-1994. As previously discussed the VG model has an integrated density and therefore the MLE could be performed without any filtering.

The statistical (historical) parameters are

\[ (\sigma = 0.117200, \theta = 0.0056, \nu = 0.002) \]

While their risk-neutral parameters are

\[ (\sigma = 0.1213, \theta = -0.1436, \nu = 0.1686) \]
Again we can see that the historical estimate for $\theta$ is close to zero, while the risk-neutral one is significantly negative. This negative $\theta$ is what creates the negative skewness observed in cross-sectional estimations.

We can try to reproduce the above parameters with the VGSA model. The resulting time-series parameter-set is

$$(\kappa = 79.499687, \eta = 3.557702, \lambda = 0.000000)$$

$$(\sigma = 0.049656, \theta = 0.006801, \nu = 0.008660, \mu = 0.030699)$$

Although the results seem to be very different, upon simulation we can see that even if the resulting arrival-rates and Gamma variables are different, the Log stock prices are close. This can be seen in Figures 3.20, 3.21 and 3.22.

An alternative would be to use the EPF algorithm with the VGSA model over the same period, in which case we would obtain

$$(\kappa = 190.409721, \eta = 3.459288, \lambda = 5.430759)$$

$$(\sigma = 0.050243, \theta = 0.002366, \nu = 0.007945, \mu = 0.032576)$$

Once again the most unstable parameters are $(\kappa, \eta, \lambda)$ or the ones corresponding to the arrival-rate. We have seen this many times, the estimation of the parameters affecting the noise is less reliable. This is in agreement with what we had observed in the second chapter, and shows the limitations of these inference tools.
Figure 3.20: Arrival Rates for simulated SPX prices using $\Psi = (\kappa = 0.0000, \eta = 0.0000, \lambda = 0.000000, \sigma = 0.117200, \theta = 0.0056, \nu = 0.002)$ and $\Psi = (\kappa = 79.499687, \eta = 3.557702, \lambda = 0.000000, \sigma = 0.049656, \theta = 0.006801, \nu = 0.008660, \mu = 0.030699)$. We can see that they are quite different.
Figure 3.21: Gamma times for simulated SPX prices using $\Psi = (\kappa = 0.0000, \eta = 0.0000, \lambda = 0.000000, \sigma = 0.117200, \theta = 0.0056, \nu = 0.002)$ and $\Psi = (\kappa = 79.499687, \eta = 3.557702, \lambda = 0.000000, \sigma = 0.049656, \theta = 0.006801, \nu = 0.008660, \mu = 0.030699)$. 
Figure 3.22: Log stock Prices for simulated SPX prices using $\Psi = (\kappa = 0.0000, \eta = 0.0000, \lambda = 0.000000, \sigma = 0.117200, \theta = 0.0056, \nu = 0.002)$ and $\Psi = (\kappa = 79.499687, \eta = 3.557702, \lambda = 0.000000, \sigma = 0.049656, \theta = 0.006801, \nu = 0.008660, \mu = 0.030699)$. Unlike arrival rates, the spot prices are hard to distinguish. This is consistent with our previous observations.
Cross-Sectional vs. Time-Series VGSA

Applying the Particle Filtering algorithm described in the previous chapter to S&P500 we find for 2001-2003 the statistical parameter-set

\[
\begin{align*}
&\kappa = 55.01778, \eta = 3.721583, \lambda = 8.666717, \sigma = 0.118637, \theta = 0.060053, \nu = 0.00103 \\
&\mu_S = -0.2910
\end{align*}
\]

and for the 1995-1999 time-period

\[
\begin{align*}
&\kappa = 1.151952, \eta = 5.418226, \lambda = 2.840461, \sigma = 0.055811, \theta = 0.008626, \nu = 0.006021 \\
&\mu_S = 0.249051
\end{align*}
\]

A typical Cross-Sectional risk-neutral parameter-set

\[
\begin{align*}
&\kappa = 2.72, \eta = 2.18, \lambda = 5.68, \sigma = 0.21, \theta = -0.41, \nu = 0.06
\end{align*}
\]

As we can see the implied skew and kurtosis are stronger for the cross-sectional method comparing to the statistical one. This is consistent with results observed with other diffusion-based models.

We perform more recent parameter-estimations corresponding to 19990610-20030610 and 19990910-20030910 (via PF based upon 1000 particles) for S&P500. The results are reported in table 3.3. As we can see, the algorithm for the estimation of the statistical parameters seems fairly stable provided that the initial parameters are chosen sufficiently close to the optimal ones.

The cross-sectional results could be computed the same way as for diffusion based models. Quoting the results of Carr et al. [48] we have the table 3.4. As we can see, for some periods the risk-neutral implied \( \lambda \) is much larger than the statistical one. This implies the possibility of a Skewness Trade as previously discussed.

<table>
<thead>
<tr>
<th>period</th>
<th>( \kappa )</th>
<th>( \eta )</th>
<th>( \lambda )</th>
<th>( \sigma )</th>
<th>( \theta )</th>
<th>( \nu )</th>
</tr>
</thead>
<tbody>
<tr>
<td>990910-030910</td>
<td>5.131967</td>
<td>6.499669</td>
<td><strong>4.360002</strong></td>
<td>0.087000</td>
<td>-0.024862</td>
<td>0.002000</td>
</tr>
<tr>
<td>990610-030610</td>
<td>6.514068</td>
<td>6.500001</td>
<td><strong>4.360000</strong></td>
<td>0.085000</td>
<td>-0.025000</td>
<td>0.001800</td>
</tr>
</tbody>
</table>

Table 3.3: VGSA statistical parameters estimated via PF. The stock-drifts \( \mu_S \) are \( -0.009999 \) and \( -0.010000 \) respectively.

It therefore seems that depending on the period, the statistical and risk-neutral parameters \( \lambda \) may or may not be consistent.
3.6 A Word of Caution

Accuracy issues of the Inference tools aside, there are practical considerations we need to bear in mind. We are applying basic models such as Heston or VGSA to a complex and constantly changing market. Indeed the true dynamics of the stock and option markets are unknown, and even if the above models approximate them fairly well, there is no guarantee that there will not be a mutation in future dynamics. The best we can do is to use the information hitherto available and hope that the future behavior of the assets is not too different from the past one. Needless to say, as time passes by and new information becomes available, we need to update our models and parameter values. This could be done within either a Bayesian or Classical framework. Therefore detecting an inconsistency between the stock and option markets does not allow us to make a riskless profit, since we simply do not know what the future is reserving for us. Once again the Skewness Transaction described in this chapter is more similar to Selling Insurance than an Arbitrage.
Bibliography


The Volatility Process


[67] Cox J. C., Rubinstein M. (1985) “Options Markets” *Prentice Hall*


[84] Duan J. C. (1996) “A Unified Theory of Option Pricing under Stochastic Volatility: from GARCH to Diffusion” Hong Kong University of Science and Technology


[133] Haykin S. (2001) “Kalman Filtering and Neural Networks” Wiley InterScience


[140] Hobson D. G. (1996) “Stochastic Volatility” *School of Mathematical Sciences, University of Bath*


[219] Silva A. C., Yakovenko V.M. (2002) “Comparison between the Probability Distribution of Returns in the Heston model and Empirical data for Stock Indices” *Department of Physics, University of Maryland*


scented Particle Filter” Oregon Graduate Institute, Cambridge University 
and UC Berkeley

ematical Sciences, New York University

Nonlinear Estimation” Oregon Graduate Institute of Science and Technol-
ogy

the Unscented Transformation” Oregon Graduate Institute of Science and 
Technology

partment of Computer Science, University of North Carolina at Chapel Hill


tic Process Limits and their Application to Queues” AT&T Labs Research, 
Springer

of Financial Economics, 19

& Sons, Ltd.

ical Pricing and Computations” Oxford Financial Press

Volatility Models” Wilmott Associates

Modeling and Inference in Economics and Econometrics” University of 
Chicago

Scales: Determining Integrated Volatility with Noisy High-Frequency 
Data” Working Paper. Carnegie Melon University, University of Chicago 
and Princeton University

Valuing Defaultable Securities” Federal Reserve Board, Washington, DC