An Affine-Invariant Sampler for Exoplanet Fitting and Discovery in Radial Velocity Data

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Received 2011 April 12; accepted 2011 November 29; published 2012 January 17

Abstract

Markov chain Monte Carlo (MCMC) proves to be powerful for Bayesian inference and in particular for exoplanet radial velocity fitting because MCMC provides more statistical information and makes better use of data than common approaches like chi-square fitting. However, the nonlinear density functions encountered in these problems can make MCMC time-consuming. In this paper, we apply an ensemble sampler respecting affine invariance to orbital parameter extraction from radial velocity data. This new sampler has only one free parameter, and does not require much tuning for good performance, which is important for automatization. The autocorrelation time of this sampler is approximately the same for all parameters and far smaller than Metropolis–Hastings, which means it requires many fewer function calls to produce the same number of independent samples. The affine-invariant sampler speeds up MCMC by hundreds of times compared with Metropolis–Hastings in the same computing situation. This novel sampler would be ideal for projects involving large data sets such as statistical investigations of planet distribution. The biggest obstacle to ensemble samplers is the existence of multiple local optima; we present a clustering technique to deal with local optima by clustering based on the likelihood of the walkers in the ensemble. We demonstrate the effectiveness of the sampler on real radial velocity data.

Key words: methods: data analysis – methods: numerical – methods: statistical – planetary systems – stars: individual (HIP36616, HIP88048) – techniques: radial velocities

Online-only material: color figures

1. Introduction

Markov chain Monte Carlo (MCMC) has proven very helpful in astrophysics for searching, optimizing, and sampling probability distributions (Brooks et al. 2011). In most cases, astrophysicists use the Metropolis–Hastings (Gilks et al. 1996) algorithm with a carefully tuned proposal distribution. Those who have departed from Metropolis–Hastings often have done so not to improve speed but to improve the properties of the sampler for computing the Bayesian evidence integral (Gregory & Loredo 1992; Loredo 1999; Wandelt et al. 2004).

In particular, MCMC has proven to be powerful for identification of exoplanet signals in radial velocity (RV) data, and to quantify the reliability and accuracy of exoplanet parameter inferences (e.g., Ford 2005; Driscoll 2006; Ford & Gregory 2007). MCMC produces samples of the posterior distribution of the orbital parameters of each exoplanet around its target star (Gregory 2005; Ford 2006; Ford & Gregory 2007). In turn, the sampling permits trivial (approximate) posterior probability calculations and parameter marginalization to obtain the distribution of individual parameters or the joint distribution of several parameters. The standard Metropolis or Gibbs sampling approaches have several practical difficulties. One is that they have problem-specific computational parameters and possibly a large number of them, especially those highly correlated, require hand tuning (Ford 2006; Gregory 2011). Even with optimally tuned diagonal parameters, these methods can be very slow on some data sets. And of course, generally, there are local optima in the likelihood function that correspond to incorrect companion identifications, see Figure 1.

In this paper, we apply a new MCMC method (Gilks & Roberts 1994; Goodman & Weare 2010)—an ensemble sampler—to radial velocity analysis. This ensemble sampler has the property of being invariant under affine transformations (see below for precise definitions). This means that the new sampler automatically, and without hand tuning, works in the optimal linear rescaling of the problem. This results in much smaller autocorrelation times, which means that one gets the same amount of information about the posterior with fewer evaluations of the likelihood function. Our code has two other features that proved indispensable in analyzing exoplanet RV data. The first is a simulated annealing cooling schedule that we use to generate the initial sample. The second feature is a simple clustering method that identifies and removes samples from irrelevant local optima.

2. Model

Our goal is to determine what configurations of planets around a target star are consistent with given radial velocity measurements. The physical model consists of a collection of Keplerian orbits. The orbit of each planet is parameterized by five orbital parameters (Ford 2006; Ford & Gregory 2007; Gregory 2005; Ohta et al. 2005). This is consistent with the approximation that on the observational timescale of a few years, interactions between the orbits are negligible otherwise the orbits would not be stable. This leads to

\[ v_{\text{rad}}(t) = v_0 + \sum_p \Delta v_p(t) , \]

where \( v_0 \) is the systemic velocity of the star, \( \Delta v_p(t) \) are the velocity perturbations from the orbital motion of each planet, and \( t \) is time.

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where \( v_0 \) is a constant velocity offset, \( p \) labels the planet, and \( \Delta v_p(t) \) is the perturbation contributed by planet \( p \). The perturbation is given by

\[
\Delta v_p(t) = A_p [\sin (f_p + \varpi_p) + e_p \sin \varpi_p], \quad (2)
\]

where \( A \) is the velocity amplitude, \( \varpi \) is the longitude of periastron, \( e \) is the eccentricity, and \( f \) is the true anomaly, which depends on time. The amplitude \( A \) can be given by

\[
A = \frac{m_p}{m_s + m_p \sqrt{1 - e^2}}, \quad (3)
\]

where \( m_p \) is the mass of the planet, \( m_s \) is the mass of the star, \( \omega \) is the mean angular velocity \((2\pi / P, \text{where } P \text{ is the period})\), \( a \) is the distance between the star and the planet, and \( i \) is the inclination between the normal direction of the orbit and the line of sight. The true anomaly \( f \) is related to the eccentric anomaly \( E \) through

\[
\cos f = \frac{\cos E - e}{1 - e \cos E}, \quad (4)
\]

and \( E \) is related to the mean anomaly \( M \) through

\[
M = E - e \sin E, \quad (5)
\]

where \( M = \omega t + \phi, \phi \) is the phase of pericenter passage. Putting the above equations together, we get

\[
\Delta v_p(t) = A_p [\sin (\psi (\omega_p t + \phi_p) + \varpi_p) + e_p \sin \varpi_p], \quad (6)
\]

where \( \psi \) is a function determined by Equations (4) and (5).

There are five independent parameters for each planet in Equation (6), which are \( A_p, \omega_p, \phi_p, e_p, \) and \( \varpi_p \). We use an equivalent set of parameters, \( \omega, A_c = \sqrt{A} \cos \phi, A_s = \sqrt{A} \sin \phi, e_c = \sqrt{e} \cos \varpi, \) and \( e_s = \sqrt{e} \sin \varpi \). Although our MCMC method is not affected by linear changes of variables, this nonlinear change of variables did improve its performance. We believe this is because it bounds \( \phi \) and \( \varpi \) between 0 and \( 2\pi \) periodically, thus making full use of machine precision.

Another possible change of parameters is replacing \( \phi \) in \( A_c \) and \( A_s \) with \( \phi + \varpi \), leaving the others \((\omega, e_c, e_s)\) unchanged. This re-parameterization makes the joint phase uncertainty in \( \phi \) and \( \varpi \) less nonlinear because \( \phi \) and \( \varpi \) are linearly correlated, as seen in Figure 7. However, any nonlinear change of variables in parameter space introduces a Jacobian factor in the prior and likelihood. Replacing \( \phi \) in \( A_c \) and \( A_s \) with \( \phi + \varpi \) causes a complicated Jacobian factor, while for \( \omega, A_c, A_s, e_c, \) and \( e_s \), which is the re-parameterization we finally used, the Jacobian factor is just 1. For \( n \) planets, there are \( 5n \) parameters. Our model has two additional parameters. One is the reference velocity offset \( v_0 \). The other is the jitter, \( \sigma \), as described in the following paragraph. (If there are multiple observatories or instruments, there will be multiple velocity offsets and jitters, which can be easily included in the code.) Together, these form the \( M = 5n + 2 \) components of our parameter vector, \( \theta \in \mathbb{R}^M \).

The data for an estimation consist of \( N \) observations of the radial velocity of a given star. For observation \( i \), \( t_i \) (MJD in days) is the observation time, \( v_i \) (m \( s^{-1} \)) is the observed radial velocity, and \( \sigma_i \) (m \( s^{-1} \)) is the reported standard deviation of the observation error. This is the observer’s estimate of the measurement noise induced by the instrument. It does not include astrophysical noise such as stellar oscillation. We denote the data collectively as \( D = (t_1, v_1, \sigma_1, \ldots, t_N, v_N, \sigma_N) \). Our data model assumes that observed velocities are independent and Gaussian with mean \( v_\text{rad}(t_i) \) and variance \( \sigma_i^2 + s^2 \). The parameter \( s \) is the jitter referred to above, which consists of all the noises not included in the measurement noise estimation \( \sigma \). Jitter increases the posterior uncertainties of the parameters, so the parameter estimation is less optimistic and thus more conservative (Gregory 2005). The probability density to observe data \( D \), given parameters \( \theta \), therefore is given by the Gaussian
Table 1
Priors for the Model Parameters

| Parameter          | Variable    | Prior     | Mathematical Form |
|--------------------|-------------|-----------|-------------------|
| Amplitude          | $A$ (m s$^{-1}$) | Jefferys  | $\frac{(A+A_{\text{min}})^{-1}}{\ln \left( \frac{A_{\text{max}}}{A_{\text{min}}} \right)}$ |
| Period             | $P (s) = 2\pi/\omega$ | Jefferys  | $\frac{(P_{\text{max}})^{-1}}{\ln \left( \frac{P_{\text{max}}}{P_{\text{min}}} \right)}$ |
| Orbital phase      | $\phi (\text{rad})$ | Uniform   | $0 \leq \phi \leq 2\pi$ |
| Eccentricity       | $e$         | Uniform   | $0 \leq e \leq 1$ |
| Longitude of periastron | $\sigma (\text{rad})$ | Uniform   | $0 \leq \sigma \leq 2\pi$ |
| Jitter             | $s (\text{m s}^{-1})$ | Jefferys  | $\frac{(A_{\text{max}})^{-1}}{\ln \left( \frac{A_{\text{max}}}{A_{\text{min}}} \right)}$ |
| Reference velocity | $v_0 (\text{m s}^{-1})$ | Uniform   | $v_{\text{min}} \leq v_0 \leq v_{\text{max}}$ |

Notes. The hyperparameters used in the code are $A_0 = 10$ m s$^{-1}$, $s_0 = 10$ m s$^{-1}$, $v_{\text{min}} = -10000$ m s$^{-1}$, and $v_{\text{max}} = 10000$ m s$^{-1}$. $A_{\text{max}}$, $P_{\text{max}}$, and $s_{\text{max}}$ are in the normalization, so they are not actually used.

The likelihood function

$$p(D | \vec{\theta}) = (2\pi)^{N/2} \left[ \prod_{i=1}^{N} \left( \sigma_i^2 + s^2 \right)^{-1/2} \right] \times \exp \left[ -\sum_{i=1}^{N} \frac{(v_i - \nu_{\text{rad}}(i))^2}{2 \left( \sigma_i^2 + s^2 \right)} \right].$$

(7)

The parameters $\vec{\theta}$ enter into this formula through the dependence of $\nu_{\text{rad}}$ on the parameters $\vec{\theta}$. It is helpful to write this in terms of the negative log of the denormalized likelihood function

$$l(D | \vec{\theta}) = \sum_{i=1}^{N} \frac{1}{2} \left[ \frac{(v_i - \nu_{\text{rad}}(i))^2}{2 \left( \sigma_i^2 + s^2 \right)} + \log \left( \sigma_i^2 + s^2 \right) \right].$$

(8)

And the likelihood function is $p(D | \vec{\theta}) \propto e^{-l(D | \vec{\theta})}$.

We assume that the parameter vector $\vec{\theta}$ has a prior distribution in which the individual parameters are independent random variables whose distribution is given in Table 1. For demonstration purposes, we used a rather conventional non-informative prior. Planet identifications would be more reliable with a prior that better reflects our present understanding and expectations about the statistics of exoplanets. We will leave that as a topic for future work.

The prior density, $p(\vec{\theta})$, is the product of the appropriate densities over the $M = 5n + 2$ components of $\vec{\theta}$. The posterior density of $\vec{\theta}$ given the data is

$$p(\vec{\theta} | D) = \frac{p(\vec{\theta})p(D | \vec{\theta})}{p(D)}.$$  

(9)

In the present work, the normalization constant

$$p(D) = \int p(\vec{\theta})p(D | \vec{\theta}) d\vec{\theta}$$

is unknown and not used.

3. SAMPLING

Affine invariance. We use an ensemble sampling method that respects affine invariance. An affine transformation is an invertible mapping $\mathbb{R}^M \rightarrow \mathbb{R}^M$ with the form $\psi = S \vec{\theta} + b$, where $S$ is a non-singular matrix. Many Monte Carlo methods use diagonal or off-diagonal scaling matrices $S$ to improve performance. That is irrelevant for affine-invariant methods. If random variable $\vec{\theta}$ has probability density $\pi(\vec{\theta})$, then $\vec{\psi} = S \vec{\theta} + b$ has density

$$\pi_{S,b}(\vec{\psi}) = \pi_{S,b}(S \vec{\theta} + b) \propto \pi(\vec{\theta}).$$

(10)

A sampler is affine invariant if the MCMC transition probability density $P(\vec{\theta} \rightarrow \vec{\theta}') = P(\vec{\theta}' | \vec{\theta})$ transforms in the same way:

$$P(\vec{\psi} | \vec{\psi}) = C_{S,b} P(\vec{\psi}' | \vec{\theta}).$$

If $\vec{\psi}' = S \vec{\theta} + b$ and $\vec{\psi} = S \vec{\theta} + b$, where $C_{S,b}$ is a normalizing determinant independent of $\vec{\theta}$ and $\vec{\theta}'$. In our codes, this is a consequence of the fact that if the code starting with $\vec{\theta}_1$ would produce $\vec{\theta}_2, \vec{\theta}_3, \ldots$, then starting with $\vec{\psi}_1$, it would produce $\vec{\psi}_2, \vec{\psi}_3, \ldots$, always with $\vec{\psi}_j = S \vec{\theta}_j + b$.

Common MCMC samplers require certain customization to sample ill-shaped densities efficiently (Ford 2006; Gregory 2011). For the density such as the left side of Figure 7, single variable Gibbs sampler updates and isotropic Metropolis samplers require a small step size to achieve a reasonable acceptance probability. But an affine transformation can turn the highly elliptical density on the left to something more spherical, as on the right. The affine-invariant MCMC sampler does not view the ill-shaped density on the left more difficult than the well-shaped density on the right (Goodman & Weare 2010). This makes it efficient on many ill-shaped densities, without customization.

Ensemble sampling. We use an ensemble sampler that updates $L$ samples together. An ensemble, $E$, consists of $L$ samples, or walkers, $\vec{\theta}_k$. The walkers are samples of the posterior parameter distribution, $\vec{\theta}_k \in \mathbb{R}^M$. The ensemble $E = (\vec{\theta}_1, \ldots, \vec{\theta}_L)$ of $L$ walkers is a point in $\mathbb{R}^{LM}$. The target probability density for the ensemble is

$$\Pi(E) = p(\vec{\theta}_1 | D)p(\vec{\theta}_2 | D) \cdots p(\vec{\theta}_L | D).$$

(11)

That is to say that each walker is an independent sampler of the posterior.

Stretch move. In the ensemble MCMC sampler, one step of the Markov chain $E(t) \rightarrow E(t + 1)$ consists of updating all walkers one-by-one as a whole cycle. Expressed in pseudo code, it is

$$E_{|k|}(t) = (\vec{\theta}_1(t + 1), \ldots, \vec{\theta}_{k-1}(t + 1), \vec{\theta}_k(t), \ldots, \vec{\theta}_L(t)),$$

has density (11), then $E_{|k|}(t)$ has density (11) too. The update of walker $\vec{\theta}_k$ uses the current positions of the other walkers in the ensemble, which we call the complementary ensemble:

$$F_{|k|}(t) = (\vec{\theta}_1(t + 1), \ldots, \vec{\theta}_{k-1}(t + 1), \vec{\theta}_{k+1}(t), \ldots, \vec{\theta}_L(t)).$$

This is $E$ with its most current locations, but with walker $\vec{\theta}_k$ omitted.

In this work, we update $\vec{\theta}_k$ using the stretch move, defined as follows. The random stretch variable, $Z$, has density $g(z)$ described below. First, choose another walker $\vec{\theta}_j \in F_{|k|}(t)$, with
all walkers in \( F_{i0}(t) \) being equally likely. Then propose a move \( \tilde{\theta}_k \rightarrow \tilde{\psi} = Z(\tilde{\theta}_k - \tilde{\theta}_j) \). The proposed new position of \( \tilde{\theta}_k \) lies on the line containing \( \tilde{\theta}_k \) and \( \tilde{\theta}_j \), but the distance between them is stretched by \( Z \). Accept \( \tilde{\psi} \) with Metropolis probability

\[
\min\left\{ 1, Z^{M-1} \frac{p(\tilde{\psi}|D)}{p(\tilde{\theta}_k(t)|D)} \right\},
\]

where \( M \) is the dimension of the parameter space. If \( \tilde{\psi} \) is accepted, then \( \tilde{\theta}_k(t + 1) = \tilde{\psi} \). Otherwise \( \tilde{\theta}_k(t + 1) = \tilde{\theta}_k(t) \).

The density of the stretch factor, \( g(z) \), must satisfy the symmetry condition

\[
g\left(\frac{1}{z}\right) = z g(z).
\]

As long as Equation (13) is satisfied, the move \( E_{[k-1]}(t) \rightarrow E_{[k]}(t) \) satisfies detailed balance for the density (11), see Goodman & Weare (2010). A simple distribution that satisfies this condition is

\[
g(z) = \begin{cases} 
\frac{1}{z}, & z \in \left[\frac{1}{a}, a\right], \\
0, & \text{otherwise}.
\end{cases}
\]

The two parameters in the stretch move ensemble sampler are the ensemble size, \( L \), and the stretch factor parameter, \( a \). The normalization constant is \( C = \frac{1}{2} (\sqrt{a} - \frac{1}{\sqrt{a}}) \). The method requires \( L > M \) in order to sample the entire parameter space. In the runs reported here, we took \( L = 1000 \) and \( a = 2 \). The results are insensitive to ensemble size. The parameters are not tuned for individual star data sets.

In pseudo code, moving \( \tilde{\theta}(t) \rightarrow \tilde{\theta}(t + 1) \) by one stretch move is given by (Goodman & Weare 2010)

for \( k = 1, \ldots, L \)

choose \( j \in \{1, \ldots, L\} \) randomly, with \( j \neq k \)
generate \( \tilde{\psi} = \tilde{\theta} + Z(\tilde{\theta}_k - \tilde{\theta}_j), Z \) satisfying (14)
accept, set \( \tilde{\theta}_k = \tilde{\psi} \), with probability (12)
otherwise reject, leave \( \tilde{\theta}_k \) unchanged.

4. LOCAL OPTIMA

Local optima of the negative log-likelihood function (8) are a problem for many parameter estimation and sampling problems. Local optima correspond to parameter fits that are better than other nearby fits but worse than fits in different parts of parameter space. Figure 1 gives an example for one specific star data set. While any valid MCMC sampler would eventually find samples near the global optimum, doing so may require an impractically long run. With the initialization just described, after cooling (see below), walkers from the ensemble find themselves in various local probability wells. The computational challenge is twofold. We seek to explore parameter space well enough that at least some of the walkers find the well of the global optimum. We also seek to cluster walkers in the various wells so that we can remove walkers from all but the most important one, or ones.

It is clearly undesirable to manually initialize the MCMC code star-by-star to avoid local optima. To be most useful, a code should find samples near the global optimum by itself. There seems to be no practical way to guarantee this, but our combination of ensemble sampling with simulated annealing and clustering does it more reliably than our earlier codes. Also, the faster equilibration time of the ensemble sampler allows it to move more easily from local well to local well. For HIP36616, we believe our algorithm has identified the global optimum. However, in other data sets, our algorithm failed to find fits as good as those other groups found by hand using the periodogram. Further progress on this issue would be very important in creating automatic data analysis routines that do not require individual human attention for each star. For instance, right now, we only use the periodogram to roughly guess the period (see below), but we could make further use of the periodogram in our code to improve automatization.

Simulated annealing. Simulated annealing (Kirkpatrick et al. 1983) is a Monte Carlo method for finding global optima in problems that have many local optima. It uses a modified posterior distribution with an artificial “inverse temperature” parameter, \( \beta \):

\[ p_{\beta}(\hat{\theta}|D) \propto p(\hat{\theta})e^{-\beta l(\hat{\theta}|D)} , \]

where \( l \) is the negative log-likelihood function (8). Small \( \beta \) allows the sampler to explore parameter space freely without getting stuck in local wells. The desired posterior (7) corresponds to \( \beta = 1 \). The autocovariance function, described below, decays faster at small \( \beta \). Cooling (increasing \( \beta \)) slowly allows the walkers to find promising local optima. A cooling schedule is an algorithm that increases \( \beta \) during the sampling process.

We use simulated annealing in the first phase of our sampler. We initialize the ensemble as a small ball centered at a random position in parameter space except for the dimensions related to the period, for which we use the periodogram (Brethorst 1988; Cumming 2004) to check for peaks between the shortest observation interval and the total observation time to make a guess. As the data set can reveal periods outside this range, this is not completely reliable. In a one-planet fit, we use the period with the highest peak \( P_{\text{highest}} \) as the initial period guess unless the data show a linear trend, which indicates a long-period companion. In such cases we use \( P_{\text{highest}}/100 \) as the period guess. In a two-planet fit, \( P_{\text{highest}} \) is the initial guess for the first planet and for the second planet, we use the period of the second highest peak to initialize a portion of walkers and the period of the third highest peak for another portion of walkers and so on. If the data show a linear trend, we would use \( P_{\text{highest}}/100 \) as the guess for the second planet. The number 100 is chosen empirically and any reasonably large number should do. Ideally, we want our initialization to be as random as possible to demonstrate the value of the algorithm, but not as random so that the algorithm is unnecessarily sabotaged by the randomness. We initialize \( \beta \) by \( \beta_{\text{initial}} = 1/N \), where \( N \) is the size of data set so that only one datum matters on average in the beginning. The runs here used the following cooling schedule: \( \beta \) is increased in \( N - 1 \) steps and the amount is determined by \( 1/\beta_i - 1/\beta_{i+1} = 1/(N\beta_{\text{initial}}) \) where \( i \) is the index of the steps. \( \beta \) is kept constant within every one \( N \)th of the total steps. In general, we used \( N = 10 \). At the end of the cooling phase the sampler has been using \( \beta = 1 \) for a while, so the walkers in the ensemble are well equilibrated in their local optima. A typical example is the left frame of Figure 2, with several local optima clearly visible.

Clustering. Figure 1 shows the results for a particular data set after annealing, for which about 85% of the walkers in the ensemble are in a well that corresponds to an excellent fit. About 12% of the remaining walkers are in another local
The left frame contains all the walkers in the ensemble after annealing. The right frame contains only those walkers that the code identified as being in the deepest well, corresponding to the best fit.

(A color version of this figure is available in the online journal.)

well that corresponds to a worse fit (see insets). The negative log likelihood for this worse fit is an order of magnitude larger. The clustering phase of our algorithm seeks to identify clusters of walkers corresponding to the local wells, so that all but the important ones can be removed. The right frame of Figure 1 shows the result in this case. Only walkers from the global optimum remain.

While there are many sophisticated clustering methods, we found that for our problems, a simple one-dimensional clustering method is more effective and reliable than the others we tried, such as clustering in parameter space. Note that there could also be multiple local optima due to aliasing, they could have very similar likelihoods. In such cases, we would need more sophisticated clustering methods. After the annealing is finished, but before clustering, we collect the average negative log-likelihood function for each walker. (One can also use a posterior function, since the normalization constant does not matter when one is taking the difference of posterior.)

This results in the $L$ numbers

$$I_k = \frac{1}{T} \sum_{t=1}^{T} l(\tilde{\theta}_k(t)|D). \quad (16)$$

The idea is that $I_k$ is characteristic of the well walker $\tilde{\theta}_k$ is in, so that walkers in the same well will have similar $I_k$. Given $I_k$, the clustering first ranks all the walkers based on $I_k$ so that $(\tilde{\theta}_1, \tilde{\theta}_2, \ldots, \tilde{\theta}_L)$ is in the order of decreasing $I_k$, or increasing $-\log I_k$. There are big jumps in $-\log I_k$ in this sequence (see Figure 1), which are fairly easy to identify and thus to separate the jumps. We calculated the difference in $-\log I_k$ for every adjacent pair of $\tilde{\theta}_k$ starting from $k = 1$ and find the first pair whose difference is a certain amount of times bigger than the average difference before. So if

$$- \log I_{(j+1)} + \log I_{(j)} > \text{Const} \frac{- \log I_{(j)} + \log I_{(1)}}{j - 1}, \quad (17)$$

all the $\tilde{\theta}_k$ with $k > j$ are thrown away and only the ones with $k \leq j$ are kept.

A more rigorous approach to the multiple well/multiple fit problem would be to estimate the evidence integral corresponding to a well before deleting it. It could happen that a small but deep well has less posterior probability than a broad but shallow well. We do not find that this occurs in practice in the fits discussed in this paper. For example, our prior prevents very short period orbits that might give excellent but spurious fits to the data. The insets of Figure 1 suggest that the local wells should be removed in our problem. In future work we hope to address this issue more carefully by estimating the evidence integrals. Even then, clustering as we do it here will be the first step.

5. DATA AND RESULTS

We tested our code on data from the Lick K-Giant Search (Frink et al. 2002; Mitchell et al. 2003; Hekker et al. 2006, 2008; Quirrenbach et al. 2011). In this paper, we present the results for two stars: HIP36616 and HIP88048. HIP36616 is interesting because the data show a companion with a small period and small mass and another companion with very large period and very large mass. The information on the large period is incomplete, which causes difficulty for the Metropolis–Hastings algorithm in finding a good fit. HIP88048 has near-complete information on both periods so it is an easier case for Metropolis–Hastings and is used as a comparison. Other groups ran these two stars with common Metropolis–Hastings routines. They confirmed that they could easily find a good fit for HIP88048. But for HIP36616, a good fit was found only
after new data were obtained. Our code successfully found a fit for HIP36616 before the new data came in.

HIP36616 is interesting and challenging both from a physical and a sampling point of view. The RV data together with a good fit are shown in Figure 3. The data are well explained by a small close companion in a roughly circular orbit and a larger and more distant companion in a highly elliptical orbit. The mass ratio in the posterior distribution (assuming coplanar orbits) is

$$\frac{m_2}{m_1} = \frac{A_2 \sqrt{1 - e_2^2}}{A_1 \sqrt{1 - e_1^2}} \frac{\omega_1^{1/3}}{\omega_2^{1/3}} = 69.8 \pm 1.7.$$ 

The parameters with their confidence intervals are listed in Table 2. Histograms of the individual parameters are shown in Figure 4.

The second example fit with the new sampler, HIP88048, is shown in Figure 5. The parameters with their confidence intervals are listed in Table 3. We also run our code on only the first half of the data of HIP88048. We get similar results as with all the data, but with a larger variance, especially for the parameters of the planet with larger period.

6. COMPARISON WITH METROPOLIS–HASTINGS

All non-trivial MCMC samplers produce autocorrelated samples. An important measure of the effectiveness of the sampler is its covariance $C(t)$ of the equilibrium time lag $t$. The longer it takes the covariance $C(t)$ to decay to 0, the longer it takes the sampler to generate independent samples, because non-zero covariance indicates correlation between samples. To be more precise, suppose $V(\tilde{\theta})$ is some function of the parameters, such as $V(\tilde{\theta}) = \theta_j$ or some nonlinear function of the components. The equilibrium autocovariance function is

$$C_V(t) = \lim_{t_0 \to \infty} \text{cov}[V(\tilde{\theta}(t_0 + t)), V(\tilde{\theta}(t_0))].$$

The limit $t_0 \to \infty$ is only to ensure that the Monte Carlo has reached steady state. The dimensionless version of this is the
Figure 4. Histograms of the parameter posteriors from HIP36616.
(A color version of this figure is available in the online journal.)
autocorrelation function
\[
\rho_V(t) = \frac{C_V(t)}{C_V(0)} = \frac{\lim_{n \to \infty} \text{cov}[V(\bar{\theta}(t_0 + t)), V(\bar{\theta}(t_0))]}{\text{var}[V(\bar{\theta}(t_0))]} . \tag{18}
\]

We used the standard estimators
\[
\hat{C}_V(t) = \frac{1}{N - t} \sum_{n=0}^{N-t} (V(\bar{\theta}(n + t)) - \overline{V})(V(\bar{\theta}(n)) - \overline{V})
\]
and
\[
\hat{\rho}_V(t) = \frac{\hat{C}_V(t)}{\hat{C}_V(0)} .
\]

Figure 6 displays the functions \(\hat{\rho}_j(t)\), which correspond to taking \(V(\bar{\theta})\) to be the \(j\)th parameter \(\theta_j\). Note \(\rho_j(t)\) can be quite different for different components of \(\bar{\theta}\) for the same data set.

The important measure of correlation for Monte Carlo is the integrated autocorrelation time (summed, actually)
\[
\tau = \sum_{t=-\infty}^{\infty} \rho(t) = 1 + 2 \sum_{t=1}^{\infty} \rho(t) .
\]
This is the number of Monte Carlo steps needed to produce an effectively independent sample of the posterior (see Goodman & Weare 2010 and references therein).

Metropolis sampler proposal and tuning. For comparison purposes we coded a traditional MCMC sampler that updates the components \(\theta_j\) one at a time using a trial that is uniform in an interval \([\theta_j - r_j, \theta_j + r_j]\). The ranges \(r_j\) were tuned individually for each component to give acceptance probability 0.4, which is commonly thought to be roughly optimal. The tuning parameters are different for each \(j\), and for a given \(j\) they are different for each data set. One can also use a Gaussian distribution for a trial move, which should give similar results.

It is noteworthy that for the Metropolis sampler to work best, one should tune all the free parameters which are on the order of the dimension squared. But how the free parameters are tuned highly depends on where the walker is in the parameter space. That is to say, the free parameters best tuned for a place far from the best-fit optimum in the parameter space could be quite different from those best tuned for a place close to the best-fit optimum. Even if the best-tuned free parameters are similar everywhere in the parameter space, the tuning workload still increases as fast as the parameter space dimension squared.

Comparison. For the ensemble sampler, we used the ensemble mean
\[
\bar{V}(\bar{\theta}_n(t)) = \frac{1}{L} \sum_{n=1}^{L} V(\bar{\theta}_n(t)) \quad \tag{19}
\]
in place of a single value on the right-hand side of Equation (18) in the definition of \(\rho\), in other words, replacing \(V\) in Equation (18) with \(\bar{V}\) defined in Equation (19). (Note the possible confusion of notation: \(\theta_n\) is component \(n\) of the parameter vector \(\bar{\theta}\), while \(\theta_n\) is the \(n\)th parameter vector in an ensemble of \(L\) such parameter vectors.) It may seem unfair to compare one step of a single vector method to a single step of the ensemble method, given that an ensemble step requires updating \(L\) walkers rather than one. However, because the walkers in the ensemble are independent (see Equation (11)), doing \(L\) ensemble updates produces \(L\) effectively independent samples. Therefore, the ensemble sampler will be exactly as effective as the traditional one if it has the same autocorrelation function (Goodman & Weare 2010).

Two other factors should be mentioned. One is that the traditional sampler requires one likelihood evaluation per component per update, while the ensemble sampler requires one likelihood evaluation per vector update. That is, the traditional sampler uses a factor of \(M\) (the number of parameters) more likelihood evaluations per vector update. This is a serious consideration in the present application, where likelihood evaluations are the

\[\text{This does not apply to burn-in, so that the ensemble may suffer more work from the burn-in phase.}\]
most expensive part of the algorithm. To be sure, in many applications (though not ours) it is not so expensive to update the likelihood function after changing a single parameter. The other is that the ensemble method is more automatic in that it does not require component specific or data set specific tuning. All the runs reported here used parameters $a=2$ and $L=1000$.

Figure 6 shows the results for the two data sets, each of which is a star with two companions. In both cases, the traditional sampler relaxes the components of the easier companion more quickly, while the ensemble sampler (which updates all parameters together) relaxes all parameters at roughly the same rate. In both cases, the ensemble is much more effective in relaxing the parameters of the difficult companion. In the harder case, HIP36616, the relaxation for the ensemble sampler is at least an order of magnitude faster than the relaxation for the traditional sampler.

7. DISCUSSION

This paper reports progress toward the goal of making posterior sampling fast, reliable, and automatic for exoplanet radial velocity fitting. The sampler presented here performed well and automatically for all the individual star data sets we tried (about a hundred data points). Very high aspect ratio wells and spurious local wells were all handled without individual tuning using a single shell script. Future projects, such as hierarchical modeling of the exoplanet distribution (Hogg et al. 2010), can become impractical with traditional samplers if they are slow. Statistical information about exoplanets, such as the eccentricity distribution (Shen & Turner 2008; Hogg et al. 2010; Zakamska et al. 2011), the mass distribution (Howard et al. 2010), the mass–semimajor axis distribution (Schlaufman et al. 2009), the brown dwarf desert (Grether & Lineweaver 2006; Lecoute et al. 2010), and inclinations (Ho & Turner 2011), depend on rapid and reliable processing of star data sets. It is a limitation of this work that we have used uninformative priors, since we already have plenty of knowledge about exoplanets; we should infer from the collection of all exoplanets that priors improve our fitting for any individual new system. Hierarchical approaches make it possible to infer informative priors without making strong new assumptions. Hierarchical model comparison and selection is a future project, as are projects in which we tune our optimization strategy to operate rapidly and robustly on large numbers of exoplanet systems (a prerequisite for efficient hierarchical modeling).

We believe that Figure 7 at least partly explains the faster decay of correlations in the ensemble sampler. For the probability distribution in the left frame, traditional single-variable updates must have small proposal steps or suffer high rejection rates. For example, if $\phi_1$ is fixed, then $\sigma_1$ cannot move much and stay within the range of likely parameters. Samplers based on isotropic multivariate proposals must also take small proposal steps. But the ensemble sampler moves walkers along lines between themselves and other walkers, which naturally adapts the proposal steps to the geometry of the distribution.

In our comparison with the Metropolis–Hastings sampler, we did not tune the sampler to its full extent. If we did, it is reasonable to think that the performance of the Metropolis–Hastings sampler—locally where it is tuned—would be the same as our ensemble sampler. But tuning free parameters (whose
Goodman & Weare (2010) speculate that one could use half of the ensemble to build a model of the posterior. For example, methods. Another would be to make more use of the information in parallel. Our current implementations do not exploit this parallelism; more sophisticated software designed for high performance computing. Each walker in the ensemble evolves in parallel. Our current implementations do not exploit this parallelism; more sophisticated software designed for high performance hardware would be welcome.

One of the problems we have encountered is that our ensemble sampler is greatly affected by the existence of multiple local optima. If the walkers occupy different local optima, the performance computing. Each walker in the ensemble evolves in parallel. Our current implementations do not exploit this parallelism; more sophisticated software designed for high performance hardware would be welcome.

We believe that more sophisticated methods from machine learning could have a large impact on these sampling problems. One obvious application would be more sophisticated clustering methods. Another would be to make more use of the information in the ensemble to build a model of the posterior. For example, Goodman & Weare (2010) speculate that one could use half of the ensemble to build a nonlinear model of the posterior that goes.

The ensemble sampler may also be better suited for high performance computing. Each walker in the ensemble evolves in parallel. Our current implementations do not exploit this parallelism; more sophisticated software designed for high performance hardware would be welcome.

All of the software used for this project is available upon request. Included in this is a general purpose of the ensemble sampler that can be used for other sampling problems.

It is a pleasure to thank Jo Bovy, Debra Fischer, Daniel Foreman-Mackey, and Ros Skelton for useful discussions and feedback.

We want to particularly thank Andreas Quirrenbach for generously sharing data with us.

F.H. and D.W.H. were partially supported by the NSF (grants AST-0908357 and HSS-1124794), NASA (grant NNX08AJ48G), and the Alexander von Humboldt Foundation. J.G. and J.W. were partially supported by DOE grant DE-FG02-88ER25053.

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Figure 7. Relationship between $\phi$ and $\sigma$ of both companions of HIP36616. We used Gaussian kernel density estimation to estimate the joint probability distribution of $\phi_1$ and $\sigma_1$ (left) and $\phi_2$ and $\sigma_2$ (right) from a large number of samples.

(A color version of this figure is available in the online journal.)