Numerical simulations of a loop quantum cosmos: robustness of the quantum bounce and the validity of effective dynamics

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Received 17 March 2014, revised 6 April 2014
Accepted for publication 8 April 2014
Published 6 May 2014

Abstract
A key result of isotropic loop quantum cosmology is the existence of a quantum bounce which occurs when the energy density of the matter field approaches a universal maximum close to the Planck density. Though the bounce has been exhibited in various matter models, due to severe computational challenges, some important questions have so far remained unaddressed. These include the demonstration of the bounce for widely spread states, its detailed properties for the states when matter field probes regions close to the Planck volume and the reliability of the continuum effective spacetime description in general. In this manuscript we rigorously answer these questions using the Chimera numerical scheme for the isotropic spatially flat model sourced with a massless scalar field. We show that, as expected from an exactly solvable model, the quantum bounce is a generic feature of states even with a very wide spread, and for those which bounce much closer to the Planck volume. We perform a detailed analysis of the departures from the effective description and find some expected, and some surprising results. At a coarse level of description, the effective dynamics can be regarded as a good approximation to the underlying quantum dynamics unless the states correspond to small scalar field momenta, in which case they bounce closer to the Planck volume or are very widely spread. Quantifying the amount of discrepancy between the quantum and the effective dynamics, we find that the departure between them depends in a subtle and non-monotonic way on the field momentum and different fluctuations. Interestingly, the departures are generically found to be such that the effective dynamics overestimates the spacetime curvature, and underestimates the volume at the bounce.
1. Introduction

A reliable understanding of the physics of the very early universe and its initial conditions requires us to go beyond the realm of validity of Einstein’s General Relativity (GR). In the classical theory unless the matter is chosen to violate the weak energy condition, the backward evolution of a spatially flat Friedmann–Robertson–Walker (FRW) universe ends in a big bang singularity. The evolution of the FRW universe is hence past incomplete in GR. Singularity theorems of Penrose and Hawking show that the fate of other spacetimes in the presence of spatial curvature, anisotropies and inhomogeneities is similar. It has been long expected that this key limitation of GR—its breakdown when the singularities are approached—arises from assuming the validity of the classical notion of spacetime at all scales, and a quantum theory of spacetime and gravity would provide insights on the resolution of singularities. In the last decade, progress in quantization of cosmological models in loop quantum cosmology (LQC) has provided a glimpse of the new physics which may arise from the quantization of gravity using non-perturbative techniques in loop quantum gravity (LQG). For various matter models, a rigorous quantization has been performed, and a common result found in the study of all these models is the existence of a quantum bounce when the curvature of the spacetime becomes Planckian [1]. The backward evolution of an expanding branch of
the universe in LQC undergoes a smooth non-singular bounce, as opposed to the classical theory where the evolution is singular [2–4].

The underlying quantum evolution equation in LQC is a discrete equation in the geometric representation. In the homogeneous and isotropic spacetime, the setting which we are interested in, the quantum Hamiltonian constraint is a difference equation with a uniform spacing in volume [4–13]. This uniform spacing is fixed by the minimum eigenvalue of the area operator in LQG3. The origin of the discretization lies in the non-local character of the gravitational field strength of the Ashtekar–Barbero connection expressed in terms of holonomies [18]. The matter part of the Hamiltonian constraint is quantized using conventional Fock quantization, resulting in differential operators for the corresponding momenta. If the matter is chosen to be a massless scalar field $\phi$, it can act as an internal clock and unitary evolution can be studied using a Klein–Gordon like equation. The quantization procedure leads to some key differences at both kinematical as well as physical level with the Wheeler–DeWitt theory, including a universal maxima for the expectation value of the energy density $\rho_{\text{max}} \approx 0.409\rho_\text{Pl}$[19] and a zero probability for the occurrence of a big bang singularity [20]. However, the quantum difference equation of LQC is very well approximated by the differential Wheeler–DeWitt equation when spacetime curvature becomes much smaller than the Planckian value. For the massless scalar field model the curvature invariants can be completely captured by the energy density $\rho = \frac{p_\phi^2}{2V^2}$, which implies, since the momentum $p_\phi$ is a constant of motion, that the above agreement occurs when the physical volume $V$ is much larger than the Planck volume.

It is interesting to note that for suitable semi-classical states, it is possible to derive an effective Hamiltonian for LQC using a geometrical formulation of quantum mechanics [21–23]. The effective Hamiltonian provides a continuum description of the underlying quantum geometry in terms of the modified Friedmann and Raychaudhuri equations [24], and has served as a very valuable tool to gain many insights on the new physics in LQC [1]. A key result of the numerical simulations performed so far for the isotropic as well as anisotropic models in LQC, is that the effective dynamics obtained from the corresponding effective Hamiltonian is a very good approximation to the underlying quantum dynamics at all scales for initial states which are peaked at a large volume at late times [2–8, 25]. However, there are some important caveats. Firstly, it is to be emphasized that the effective Hamiltonian is derived for a specific set of states which are semi-classical and sharply peaked at a classical trajectory at late times when the spacetime curvature is small compared to the Planckian curvature. Therefore, departures from the effective theory can be expected if one considers states which are widely spread. Secondly, an assumption underlying the derivation of the effective Hamiltonian in [21–23] is that the volume is always greater than the Planck volume. This assumption is tied to the approximation of the eigenvalues of the inverse volume operator in the Hamiltonian constraint with $1/V$. It is remarkable that even at volumes as small as ten times the Planck volume, the error in this approximation is less than 1%. Note that in all the numerical investigations carried out so far, excellent agreement was found with the effective theory as, in these cases, the quantum bounce occurred at volumes far greater than the Planck volume. As an example, in the study of spatially flat isotropic models with a massless scalar field [4], the value of the bounce volume was greater than 1000 times the Planck volume. Moreover, the initial

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3 Due to ambiguities in the quantum theory, one may be tempted to motivate a quantum Hamiltonian constraint with a different discretization property, such as uniform discretization in the scale factor. However, it turns out that all but uniform discretization in volume are ruled out by mathematical consistency [14]. This is supported by factor ordering issues [15] and the stability arguments of the difference equation [16, 17].
states were always chosen to be sharply peaked with relative fluctuations much smaller than unity.

In previous works such as in [2–4], where the objective was to understand the fate of the classical singularity in the quantum theory for a universe which corresponds to a classical macroscopic universe at late times, the value of \( p_\phi \) at which the initial semi-classical state was peaked was taken to be large. Our objective, in this article, is to extend the present understanding of the quantum bounce in LQC to more extreme initial conditions (including small \( p_\phi \)) for which the volume at the bounce is of the order of a few Planck volumes, and probe the validity of the effective dynamics for a wide variety of initial conditions. Since the effective Hamiltonian approach is a very important tool to extract physics in LQC, it is pertinent to carefully understand the validity of the effective dynamics by probing regimes where we expect it to fail. This would be the case when the bounce happens close to the Planck volume and when the initial states are not be sharply peaked [23]. Such an exercise will allow a rigorous test of the robustness of the quantum bounce and effective dynamics for far more general states than the ones considered so far. If the departures from the quantum theory are found, then this would open a new avenue to compute finer corrections to the modified Friedmann dynamics and understand the robustness of various phenomenological predictions extracted in LQC. This can be potentially very useful. In particular to gain a deeper understanding of the novel results found in the pre-inflationary epoch in LQC [26] and by considering the perturbations of the effective Hamiltonian [27].

Probing the effects of quantum discreteness at geometrical scales much smaller than those considered so far, and to check the robustness of effective theory for states which are not sharply peaked is computationally very challenging. To understand the associated difficulties, let us first note that the technical challenges for both types of states are essentially the same since a state probing quantum geometry at much smaller scales than those considered so far will be widely spread. We note that the scale at which a state bounces, which can be regarded as the minimum volume of quantum geometry probed by the initial state, directly depends on the value of the scalar field momentum \( p_\phi \) on which the initial state is peaked. The effective dynamics predicts that the bounce occurs at volume \( V_b^{(\text{eff})} \approx 1.1 \, p_\phi \, V_\text{Pl} / \sqrt{G} \). If the field momentum is taken to be \( p_\phi = 1000 \sqrt{G} \hbar \), a typical value for which numerical simulations have been performed in LQC, then the universe will bounce at \( V_b^{(\text{eff})} \approx 1100 \, V_\text{Pl} \) according to the effective theory. Indeed, it turns out that for sharply peaked states the quantum bounce occurs at approximately the same value [2–4]. This bounce volume is larger than the underlying quantum discreteness in volume by approximately three orders of magnitude. Further, the errors resulting from the approximation of inverse volume modifications considered in the effective Hamiltonian at this bounce volume are smaller than \( 10^{-10} \). To probe the quantum geometry at volumes, say one order greater than the Planck volume, we expect to have to consider much smaller values of \( p_\phi \) than before. Such states are inevitably widely spread in volume. As an example, if an initial state is peaked at a very small value of \( p_\phi \), say \( p_\phi = 20 \sqrt{G} \hbar \) with \( \Delta p_\phi = 1.59 \) and peaked at \( V = 68 \, 700 \, V_\text{Pl} \), then it has relative spread in volume as \( \Delta V/V = 6.45 \). One is thus dealing with a very quantum state.

The computational cost involved in simulations with states with large relative fluctuation in volume with the techniques used so far is extremely high compared to sharply peaked states. A typical simulation for sharply peaked states, for example states which are peaked at \( p_\phi = 1300 \sqrt{G} \hbar \) and with initial relative volume dispersion \( \Delta V/V \approx 0.01 \), requires \( \sim 30 \, 000 \) grid points in the spatial direction. Such a simulation takes approximately 240 s on a 2.4 GHz Sandybridge computer with 16 cores. On the other hand, simulations with small \( p_\phi \) such as \( 20 \sqrt{G} \hbar \) with \( \Delta V/V \approx 6 \), requires the outer boundary in the numerical grid for volume to be placed at \( V_{\text{outer}} \approx 2 \times 10^{19} \, V_\text{Pl} \). This would, in turn, require \( 1.25 \times 10^{14} \) times more grid
points in the spatial direction. Due to the stability requirements for the numerical simulation\(^4\), this amounts to requiring a $1.25 \times 10^{14}$ times smaller time step for a stable evolution. On a similar workstation, this simulation would take approximately $10^{23}$ years if it is assumed that the simulation would fit in memory.

To tackle this challenge, we recently developed a hybrid numerical scheme, the Chimera scheme, by utilizing the fact that the LQC difference equation tends to the Wheeler–DeWitt equation in the large column regime [29]. The Chimera scheme uses a hybrid spatial grid having two components: an inner grid where evolution is performed using the LQC difference equation, and an outer grid chosen at very small spacetime curvature where evolution is performed using a discretized version of the Wheeler–DeWitt equation. Since we are free to choose the underlying discretization on the outer grid, it leads to a significant computational cost reduction. As an example, using the Chimera scheme for $p_\phi^* = 20 \sqrt{G\hbar}$ and $\Delta V/V \approx 6$, the numerical simulation can be performed in less than 10 min!

In this work, we use the Chimera scheme to perform simulations for various values of $p_\phi$ in the range between 20 to $1500 \sqrt{G\hbar}$ for a large range of peakedness properties of the initial states. We perform numerical simulations with initial states constructed via three different methods originally proposed in [3, 4]\(^5\). The quantum bounce is shown to occur for all three kinds of states and for all initial data parameters. This is an expected result from the analytical studies on genericness of the bounce in [19]\(^6\). The relative fluctuations across the bounce remain tightly constrained throughout the evolution, satisfying triangle inequalities found in the analysis of [30]. We obtain new and unexpected results when we analyze the departures of the effective theory from the underlying quantum evolution. The effective theory shows departures under certain conditions, and these departures have different features in the different methods for constructing the initial state. These differences, which are summarized in table 2, become significantly less pronounced for sharply peaked states for large values of $p_\phi$. The general expectation that the differences between the effective and the LQC theory grow as $p_\phi$ decreases and the fluctuations in volume increase, is found not to be true in general. We find that the results depend on the method chosen for the initial state construction and also on the way an initial state departs from a minimum uncertainty state with equal relative dispersions in the volume and the field momentum. For all the three methods, it is possible that the agreement between the effective theory and the quantum evolution improves if the relative fluctuation in volume is increased while keeping $p_\phi$ the same. On the other hand, if the relative fluctuation in volume is fixed, then a decrease in the value of $p_\phi$ at which the state is peaked always increases the departure between effective theory and quantum evolution for two of the three methods. For the third method, it depends on whether the initial state has absolute fluctuation in $p_\phi$ greater than a certain value.

Our results show that for an initial state peaked at a macroscopic universe at late times, the effective theory provides an excellent description of the quantum dynamics. It is only for the initial states which have large fluctuations that the large departures from the effective theory are found. Under certain conditions, there can be significant departures between the effective theory and the quantum evolution if the initial state is peaked at a small value of $p_\phi$.

\(4\) For the numerical simulations, the differential operators in the matter quantum Hamiltonian constraint need to be discretized. Unlike the discretization in volume, the numerical discretization in $\phi$ is not fixed by the underlying theory, but is constrained by the Courant–Friedrichs–Lewi (CFL) criteria for a stable evolution of the difference equations [28]. Due to the nature of the equations, any increase in the location of the spatial boundary requires a corresponding decrease in the time step for a stable evolution.

\(5\) Due to issues associated with the construction of the initial state, for the first method the smallest value of $p_\phi$ at which the initial state is peaked is chosen to be $500 \sqrt{G\hbar}$.

\(6\) Here we note that the quantum constraint considered in the exactly solvable model in [19] was different from the one considered here.
As mentioned earlier, such states have wide spread even initially and they do not correspond to a universe which evolves to a classical macroscopic universe. It turns out that the quantitative differences can be fitted by simple polynomial functions when plotted against the spread in field momentum $\Delta p_{\phi}$. Also, the departures in the large $p_{\phi}$ regime, though small, show interesting behavior with respect to the relative spread in $V$. A general result, found from the study of all the methods, is that the effective theory overestimates the value of spacetime curvature at the bounce and underestimates the bounce volume. Another general result is the non-monotonic behavior of the relative volume dispersion, as the dispersion in the field momentum varies for a fixed value of $p_{\phi}$. This result is in agreement with analytical studies performed with a different kind of initial states and a different quantum constraint [31]. The non-monotonicity in the volume dispersion is also reflected in the quantitative study of the differences between the effective and LQC trajectories, as the spread of the initial state is varied.

This manuscript is organized as follows. For completeness, we provide a brief overview of the main results we need in studying the loop quantization of homogeneous and isotropic spacetime with a massless scalar field, in section 2. Our discussion is based on [4], which we refer the reader to for details. In section 3 we briefly discuss the effective theory and modified Friedmann equation for flat FRW model. In section 4 we describe the three different methods of specifying the initial states, which were originally discussed in [3]. Here we also discuss properties of uncertainty products and minimum uncertainty states. In section 5 we briefly discuss the Chimera scheme, which is used for the numerical simulations discussed in this paper. In section 6 we show some representative results with one of the methods to provide a snap shot of the new findings. Here we show the contrast in results for sharply peaked initial states with a large value of $p_{\phi}$ at which the state is peaked, with a widely spread initial state chosen to be peaked at a small value of $p_{\phi}$. In the latter case, we show departures of the effective trajectory from the quantum expectation values of the volume observable (and their agreement within the variances) for different values of relative fluctuations. We also demonstrate in this section, the way the relative fluctuations change and the way triangle inequalities between them are satisfied for different methods of construction of initial states. In section 7, we study the differences between the effective and the LQC trajectories quantitively in detail for different initial state methods. This brings out several interesting features both in the small and large $p_{\phi}$ regime. In section 8 we provide a summary and discussion of the main results presented in the paper.

### 2. Loop quantum cosmology: spatially flat, isotropic model

Loop quantization of the homogeneous and isotropic model is a constraint based quantization which closely follows the procedure in LQG—a candidate theory for canonical quantization of gravity based on Ashtekar variables: the SU(2) connection $A^a_i$ and the conjugate triad $E^i_a$. Due to the underlying symmetries of the homogeneous and isotropic spacetime, the Ashtekar–Barbero connection and the triad can be symmetry reduced to connection $c$ and triad $p$ [18]:

$$A^i_a = c V^{1/3} \tilde{\omega}_a^i, \quad E^a_i = \text{sgn}(p) p V^{-2/3} \sqrt{q} \tilde{e}_a^i, \quad (2.1)$$

where $\tilde{e}_a^i$ and $\tilde{\omega}_a^i$ respectively denote the densitized triads and co-triads compatible with the fiducial metric $\tilde{q}_{ab}$ on the spatial manifold which we consider to be a three-torus. In the above equation, $V$ denotes the fiducial volume: $V = \int_{T^3} d^3 \sqrt{q}$. The fiducial metric is related to

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7 Results of this analysis, are independent of the particular topology of the spatial manifold.
the spatial metric $q_{ab}$ as: $q_{ab} = a^2 \delta_{ab}$, where $a$ denotes the scale factor in the spacetime line element:

$$\text{d}x^2 = -N^2 \text{d}t^2 + a^2(t)(\text{d}x^2 + \text{d}y^2 + \text{d}z^2).$$

(2.2)

In our analysis, we will choose the lapse $N = 1$. For this choice of lapse, the connection and its conjugate triad are related to the metric and its time derivative as

$$p = \dot{V} 1/3a^2, \quad c = \gamma \dot{V} 1/3\dot{a},$$

(2.3)

where $\gamma \approx 0.2375$ is the Barbero–Immirizi parameter [33, 34].

In the loop quantization of cosmological spacetimes, due to symmetry reduction, the only non-trivial constraint is the Hamiltonian constraint. Following LQG, the gravitational part of the classical Hamiltonian constraint is expressed in terms of the holonomies and fluxes of the symmetry reduced Ashtekar–Barbero connection $c$ and the conjugate triad $p$

$$C_{\text{grav}} = -\frac{1}{\gamma^2} \int d^3x \epsilon_{ijk} F^u E^{ij} F_k \frac{E^u}{\sqrt{|\text{det}(E)|}} = -\frac{6}{\gamma^2} c^3 \sqrt{|p|},$$

(2.4)

where $F_{\alpha \beta}$ denotes the field strength of the connection in terms of the holonomies over straight edges $\lambda$ $e^{\alpha}_k$ $h^{(\lambda)}_k = \cos(\lambda c/2)I + 2 \sin(\lambda c/2)\tau_k,$

(2.5)

where $\tau_k = -i\sigma_k/2$, and $\sigma_k$ denote Pauli spin matrices. The field strength $F_{\alpha \beta}$ is written in terms of the holonomy around a square loop, as in the classical gauge theory, as:

$$F_{\alpha \beta} = 2 \lim_{\Delta \rightarrow 0} \text{Tr} \left( \frac{h^{(\lambda)}_{\alpha \beta} - 1}{\lambda^2 \sqrt{V} 2/3} \right) \omega^i \dot{\omega}^j,$$

(2.6)

where

$$h^{(\lambda)}_{\alpha \beta} = h^{(\lambda)}_i h^{(\lambda)}_j \left( h^{(\lambda)}_i \right)^{-1} \left( h^{(\lambda)}_j \right)^{-1}. $$

(2.7)

In the quantum theory, due to the underlying quantum geometry, the area of the loop $\lambda^2 p$ has a non-zero minimum $\Delta = 4\sqrt{3}(\gamma\pi)^{3/2}$. Thus, $\lambda = \sqrt{\Delta}/|p|^{1/2}$. The dependence of the $\lambda$ on the triad $p$ leads to the volume representation as a natural choice of the geometric representation in the quantum theory [4, 19]. On the eigenkets $|v\rangle$ of the volume operator $\hat{V} = \frac{|p|}{3/2}$, its action on the holonomy operators is of a simple translation

$$\hat{V} |v\rangle = \left( \frac{\pi \gamma}{6} \right)^{3/2} |v\rangle K |v\rangle |v\rangle, $$

(2.8)

(2.9)

The eigenkets $|v\rangle$ satisfy: $\langle v_1 | v_2 \rangle = \delta_{v_1, v_2}$, and form an orthonormal basis in the kinematical Hilbert space which is the space of the square integrable functions on the Bohr compactification of the real line.

Here we are interested in a flat FRW model with the matter part of the Hamiltonian constraint provided by a massless scalar field $\phi$: $C_\phi = 8\pi G |p|^{3/2} p^2/4 \phi$. The physical states, $\hat{\Psi}(v, \phi)$, are obtained by solving ($\hat{C}^{grav}_\phi + \hat{C}^-_\phi$) $\Psi = 0$, which yields a Klein–Gordon type equation with $\phi$ playing the role of internal time [4]:

$$\frac{\partial^2}{\partial \phi^2} \hat{\Psi}(v, \phi) = -\hat{\Theta} \hat{\Psi}(v, \phi).$$

(2.10)

$^8$ The following relation between $c$ and the time derivative of the scale factor is true only in the classical theory.
Here $\Theta$ is the discrete spatial Laplacian defined as,
\begin{equation}
\hat{\Theta} := -\frac{1}{B(v)} [C^+(v)\Psi(v+4, \phi) + C^0(v)\Psi(v, \phi) + C^-(v)\Psi(v-4, \phi)]
\end{equation}
where the coefficients $C^+$ and $C^-$ are given as
\begin{align*}
C^+(v) &= \frac{3\pi KG}{8} |v+2||v+1|-|v+3|, \\
C^-(v) &= C^+(v-4) = \frac{3\pi KG}{8} |v-2||v-3|-|v-1|, \\
C^0(v) &= -C^+(v) - C^-(v).
\end{align*}
and $B(v) = 8\frac{K}{\gamma l^3} (|[v+1]|^{1/3}-|[v-1]|^{1/3})^3$ denotes the eigenvalue of the inverse volume operator,
\begin{equation}
|\rho|^{-3/2}|v| = \left(\frac{8\pi \gamma l^3}{6}\right)^{-3/2} B(v)|v|.
\end{equation}
The LQC evolution operator given in equation (2.11) is a difference operator with a uniform discretization in $v$. As a result, the evolution equation in LQC turns out to be a difference equation, as opposed to Wheeler–DeWitt theory, where the evolution equation is a partial differential equation.

Since we are considering non-fermionic matter, physics does not distinguish between the orientations of the triads. The physical states are therefore chosen to be symmetric under the change of sign of the physical triads: $\Psi(v, \phi) = \Psi(-v, \phi)$. The inner product of the physical states can be obtained by using a group averaging procedure \cite{35, 36}
\begin{equation}
\langle \Psi_1 | \Psi_2 \rangle = \sum_{v} \overline{\Psi_1(v, \phi)} B(v)^{-1} \Psi_2(v, \phi).
\end{equation}
To extract physical predictions, we need a set of Dirac observables. As the field momentum is a constant of motion in the classical theory, it turns out to be one of the Dirac observables.

Another Dirac observable is the expectation value of the field $\phi$ at a fixed ‘emergent time’ $\phi_o, v|\phi=\phi_o$. The expectation value of these Dirac observables are computed for each value of the scalar field $\phi$
\begin{align}
\langle \Psi | \hat{\phi}|_{\phi=\phi_o} |\Psi \rangle &= |\Psi||^{-1} \sum_v B(v)|v| \Psi(v, \phi_o)|^2 \\
\langle \Psi | \hat{\phi}|^2 |\Psi \rangle &= |\Psi||^{-1} (-i\hbar) \sum_v B(v) \Psi(v, \phi) \partial_{\phi} \Psi(v, \phi).
\end{align}
Similarly, one can obtain the fluctuations of the expectation values. Note that the physical states have support on the discrete lattices $\pm \epsilon + 4n$ where $\epsilon \in [0, 4)$. These are preserved by the dynamical evolution, and hence there is a superselection in the physical Hilbert space. We will focus our analysis on the case of the lattice defined by $\epsilon = 0$ which includes the possibility of physical volume becoming zero in the evolution.

With the quantum evolution equations available, we can now consider a suitable initial state and study its evolution. By computing the expectation values of the Dirac observable, and comparing them with the classical trajectory, physical implications of the underlying quantum geometry can be studied. For the model under consideration, this task was first completed in [2–4], for initial states, with reasonably large $p_\phi$, which are sharply peaked on a classical trajectory at late times and at large volumes. The backward evolution of such states revealed the existence of a quantum bounce in the Planck regime, when the energy density of the scalar field reached a universal maximum $p_{\text{max}} \approx 0.409 p_P$ \cite{19}. It also turns out that the quantum evolution is very well approximated by a trajectory obtained from an effective Hamiltonian (see section 3) at all scales, including the bounce for sharply peaked states.
As discussed in section 1, our goal in this manuscript is to study the above quantum evolution in more detail to understand the way the quantum bounce and the agreement with the effective dynamical trajectory, obtained from the effective Hamiltonian, are affected by considering states which may not be sharply peaked and correspond to small values of $p_\phi$. For this purpose, and to facilitate a comparison with previous works, we will consider the form of the initial states identical to the ones in [2–4] in section 4. These initial states are constructed using an important property of the LQC quantum difference equation that in the large volume approximation, it is extremely well approximated by the Wheeler–DeWitt equation. In terms of $v$, the Wheeler–DeWitt equation can be written as [4]

$$\frac{\partial^2}{\partial \phi^2} \Psi(v, \phi) = -\hat{\Theta} \Psi(v, \phi) := 12\pi G v \frac{\partial}{\partial v} \left( v \frac{\partial}{\partial v} \right) \Psi(v, \phi),$$

(2.16)

where $\hat{\Theta}$, unlike $\hat{\Theta}$ in equation (2.11), is a differential operator. It is important to note that the underlying geometry in the Wheeler–DeWitt theory is continuous unlike the discrete quantum geometry in LQC. It is only in the regime where the spacetime curvature is small, that the discreteness of quantum geometry is well approximated by the smooth continuum description.

3. Effective dynamics

Before we proceed to the construction of the initial states, we briefly discuss the main features of the effective theory. The effective spacetime description is derived using the geometric formulation of quantum mechanics. Using the inner product in the Hilbert space, one can define a Riemannian metric and the symplectic structure, and treat the Hilbert space as a quantum phase space. In order to obtain the modifications to the classical dynamical equations resulting from the quantum theory, one then relates the quantum phase space variables to their classical counterparts [37]. In LQC, this has been achieved using two independent approaches under different sets of assumptions. The first is the embedding approach [21, 22] which has been extensively used for phenomenological applications and analysis of cosmological perturbations in LQC, and which is also used in this manuscript. The second is the truncation approach [38] which relates the expectation values of the basic conjugate variables in the quantum phase space to those in the classical phase space using an order by order approximation.

The embedding approach is based on utilizing a fiber bundle structure in the quantum phase space whose base space is the classical phase space. Using the expectation values of the basic quantum phase space variables, which remain constant along the fibers of the bundle, one seeks a projection of the quantum phase space into the classical phase space. In particular, with a judicious choice of states, one aims to construct approximately horizontal sections which are preserved under evolution. If such sections can be found, then a faithful embedding of quantum phase in the classical phase space is achieved and corrections to the classical dynamical equations can be obtained. In LQC, this approach has been used for Gaussian states to compute modifications to the classical Friedmann equation in the spatially flat model with a massless scalar field [22], and for dust and radiation matter fields albeit for an older form of quantum constraint [21]. These results have been extended to a general equation of state of matter recently [23]. The validity of the modified Friedmann dynamics in the effective theory has so far been verified with the evolution obtained from the quantum Hamiltonian constraint in LQC, using sharply peaked states, for isotropic spacetimes in the presence and absence of spatial curvature for massless scalar field [4, 5, 8], in the presence of a cosmological constant [6, 7] and inflationary potential [13], as well as in the case of vacuum Bianchi-I spacetime [39]. Recently, these tests have also been conducted in the case of a cyclic potential [40].
In the truncation approach, which in principle is applicable for a wider variety of states, there are infinitely many moments which are coupled by nonlinear differential equations. By taking a truncation to a lower order of moments, one can compute corrections (valid to that small order) to the classical dynamical equations. In this approach, it is important to check the self-consistency of the truncated equations and to also ensure that ignored higher moments remain insignificant throughout the evolution. Otherwise the truncated equations may not reliably capture the full quantum evolution from the quantum Hamiltonian constraint.

In the LQC literature, the phenomenological applications of the truncation approach are not as widely studied as the embedding approach and few tests have been performed to determine how well the truncation approach agrees with the full quantum evolution. There have been numerical investigations of the backreaction effects due to higher order moments on Gaussian initial states in this approach to the effective theory, see [41]. In the following, we will focus our discussion to the effective dynamics as derived using the embedding approach.

In the embedding approach, for the case of the massless scalar field in the spatially flat isotropic model the effective Hamiltonian constraint up to the terms which depend on the fluctuations is given as:

$$H_{\text{eff}} = -\frac{3V}{8\pi G\gamma^2} \frac{\sin^2(\lambda \beta)}{\lambda^2} + H_\phi \approx 0,$$

where

$$H_\phi = \frac{p^2}{2V}. \quad (3.1)$$

Here $\beta$ is the conjugate variable to $V$, satisfying $[\beta, V] = 4\pi G\gamma$. In the derivation of the above constraint, the initial state is peaked at small spacetime curvature at a large volume. The constraint is derived under the approximation that volume remains larger than the Planck volume$^9$ and the relative fluctuations of the initial state remain small compared to unity throughout the evolution [23].

Using Hamilton’s equation, the time derivative of volume can be easily computed:

$$\dot{V} = -4\pi G\gamma \frac{\partial}{\partial \beta} H_{\text{eff}} = \frac{3}{2\gamma \lambda} \sin(2\lambda \beta)V, \quad (3.2)$$

where $\lambda$ is the same value as in the quantum theory: $\lambda = 2(\sqrt{3\pi \gamma})^{1/2}l_{\text{Pl}}$. Using the effective Hamiltonian constraint given in equation (3.1), and the expression for energy density $\rho = p^2/(2V^2)$ we obtain the following modified Friedmann equation

$$H^2 = \left( \frac{\dot{V}}{V} \right)^2 = \frac{8\pi G}{3} \rho \left( 1 - \frac{\rho}{\rho_{\text{max}}^{(\text{eff})}} \right), \quad (3.3)$$

where $H$ is the Hubble rate and $\rho_{\text{max}}^{(\text{eff})}$ is the upper bound on the energy density given as

$$\rho_{\text{max}}^{(\text{eff})} = \frac{3}{8\pi G\gamma^2\lambda^2} \approx 0.409\rho_{\text{Pl}}. \quad (3.4)$$

The modifications to the classical Friedmann equation, that are independent of the choice of state, present a very good approximation to the underlying quantum geometry, and the resulting trajectories agree very well with the full quantum evolution as long as the volume is larger than the Planck volume [23]. It has also been argued that the potential corrections due to the state dependent quantum fluctuations in the non-compact model are negligible, in the limit of taking the volume of the fiducial cell to infinity [42].

Hamilton’s equation of motion for the matter field yields the following equation, which is equivalent to the conservation of the energy momentum tensor,

$$\dot{\rho} + 3H(\rho + P) = 0. \quad (3.5)$$

$^9$ As explained in section 1, the volume does not need to be exceedingly large than the Planck volume for the approximation to be valid.
Here, \( \rho \) is the energy density and \( P = -\partial H/\partial V \) is the pressure of the matter field. Using the modified Friedmann equation and the expression for the energy density of the massless scalar field, it is straightforward to find that the volume at the bounce is determined by the values of \( p_\phi \) as follows:

\[
V_b^{(\text{eff})} = 1.1051 p_\phi / \sqrt{G} \hbar. \tag{3.6}
\]

Thus, the effective dynamics predicts that the bounce volume should decrease as \( p_\phi \) is decreased. This insight plays an important role in performing numerical simulations in this paper. Later we will use the above value of \( V_b^{(\text{eff})} \) to define a quantity, \( \delta \) which quantifies the departure between the effective theory and the quantum evolution in LQC as follows:

\[
\delta = \frac{(V_b - V_b^{(\text{eff})})}{V_b^{(\text{eff})}}. \tag{3.7}
\]

4. Initial data

In the previous section we discussed the loop quantization of spatially flat isotropic FRW spacetimes. We also discussed that in the large volume limit the LQC evolution equation can be approximated by the Wheeler–DeWitt evolution equation. In this section we will utilize this property of the LQC evolution equation to construct the initial state for the numerical evolution. In all the simulations, the initial conditions are provided so that the initial energy density is very small and the initial state is peaked at a classical trajectory. Therefore the initial state can be described by a Wheeler–DeWitt state, as the Wheeler–DeWitt equation is the continuum limit of the LQC difference equation. A general solution of the Wheeler–DeWitt equation (2.16) is

\[
\Psi(v, \phi) = \int_{-\infty}^{+\infty} \Psi(\omega) \xi_\omega(v) e^{i\omega(\phi - \phi_o)} \, d\omega, \tag{4.1}
\]

where \( \Psi(\omega) \) is the wave profile in \( \omega \) space, and \( \xi_\omega(v) \) are the eigenfunction of the \( \hat{\Theta} \) operator (2.16):

\[
\xi_\omega(v) = \frac{1}{2\pi} e^{i\omega \ln(v)/\sqrt{12\pi G}}. \tag{4.2}
\]

Given a form of the wave profile, one can evaluate the above integral (numerically or analytically where possible) to obtain the form of the solution \( \Psi(v, \phi) \) at a given value of the ‘emergent time’ \( \phi \) and construct an initial state for the evolution in LQC.

In this paper, we construct the initial data in form of a Gaussian wavepacket. Such a class of initial state is characterized by three main parameters: (i) the volume at which the initial state is peaked \( v^* \) (ii) the scalar field momentum \( p_\phi \) and (iii) the spread of the state \( \sigma \) or \( \sigma_v \). The value of \( v^* \) is kept large enough to make sure the classicality of the initial trajectory. Here we will consider three different types of initial data, based on [3]. The first is a Gaussian in \( v \), and the second and third types of initial states are Gaussian in \( \omega \), characterized by \( \Psi(\omega) \) as follows

\[
\Psi(\omega) = e^{-(\omega - \omega^*)^2/2\sigma^2}, \tag{4.3}
\]

where \( \omega = p_\phi / \hbar, \omega^* \) is the value of \( \omega \) at which the initial state is peaked and \( \sigma \) represents the spread of the Gaussian waveform. The value of \( \omega^* (= p^*_\phi / \hbar) \) governs the value of the bounce volume. In the effective theory the bounce volume can be exactly predicted to be \( v_b \approx 0.32 p^*_\phi / \sqrt{G} \hbar \) [4]. Thus, larger \( p^*_\phi \) corresponds to larger volume at the bounce. The relative spread in volume \( \Delta v / v \) is inversely proportional to the spread in the field.
momentum $\sigma$. Hence, by changing $\sigma$ in the initial data, we have a control over the relative spread in volume, of the initial state.

We now discuss the three types of initial states considered in this paper.

4.1. Method-1: Gaussian in volume ($v$)

In this initial data construction method, we choose the wavefunction to be a Gaussian in $v$, peaked at large volume $v^*$. Such a state at a given value of the scalar field $\phi = \phi_o$ can be written as

$$\Psi(\phi, v) = e^{-\frac{(v-v^*)^2}{2\sigma^2}} e^{ib^*(v-v^*)}. \quad (4.4)$$

where $b$ is the variable conjugate to $v$ and $\sigma_v$ is the spread of the Gaussian waveform. Here, $b$ is related to $\beta$ in equation (3.1) as $\beta = 6 b K \sqrt{\frac{3}{16\pi G}} \gamma^2$. Using the classical Hamiltonian constraint, the $\phi$ derivative of the above wavefunction can be shown to be

$$\frac{d}{d \phi} \Psi(\phi, v) \bigg|_{\phi = \phi_o} = \sqrt{12\pi G} \frac{v(v-v^*)}{\sigma^2} + i b^* v \Psi(\phi, v). \quad (4.5)$$

where $b^*$ is the initial value of $b$, given by

$$b^* = \pm \frac{\omega^*}{\sqrt{12\pi G v^*}}. \quad (4.6)$$

Substituting the value of $b^*$ in equation (4.5) we obtain the ‘time’ derivative of the initial state and hence have the complete set of initial conditions needed in order to study the evolution. Note that the above relation between $b^*$ and $\omega^*$ holds true only in the classical limit, i.e. when the LQC difference equation can be safely approximated by the Wheeler–DeWitt equation. To satisfy this condition one has to make sure to choose $v^*$ large enough.

4.2. Method-2: Wheeler–DeWitt initial state

We consider a solution of the Wheeler–DeWitt equation given by the integral equation (4.1) as the second initial data method with the profile given by equation (4.3). Using the eigenfunctions for the Wheeler–DeWitt evolution operator (4.2), the integral in equation (4.1) yields the following form of the wave-function

$$\Psi(\phi, v) = \frac{\sigma}{\sqrt{2\pi}} \exp \left( -\frac{1}{2} \left( \frac{\ln (v/v^*) - \phi}{\sqrt{12\pi G}} - \phi \right) \left( \frac{\sigma^2 \ln (v/v^*)}{\sqrt{12\pi G}} - \sigma^2 \phi - 2i\omega^* \right) \right). \quad (4.7)$$

The initial state given above is a continuous function of $v$, but in the numerical simulations, since the spatial grid in LQC is discrete with uniform discreteness $\Delta v = 4$, we compute the states only at the discrete points on which an LQC state has support on. In order to provide all the necessary data for evolution, one also needs the value of the time derivative of state $\partial_\phi \Psi \big|_{\phi = \phi_o}$, which is given by

$$\partial_\phi \Psi(\phi, v) = \left( \frac{\sigma^2 \ln (v/v^*)}{\sqrt{12\pi G}} - \sigma^2 \phi - i\omega_o \right) \Psi(\phi, v). \quad (4.8)$$

4.3. Method-3: rotated Wheeler–DeWitt initial state

In this initial data construction method we consider a variation of method-2 described above. The initial state is obtained by rotating an initial Wheeler–DeWitt state by an $\omega$ dependent
phase factor by multiplying the eigenfunctions of the Wheeler–DeWitt operator with $e^{-i\alpha}$, where $\alpha$ is given by [3]
\[
\alpha = k(\ln(|k|) - 1)
\] (4.9)
with $k = -\omega/\sqrt{12\pi G}$. This phase is introduced to match the eigenfunctions of the LQC evolution operator with those of the Wheeler–DeWitt operator in the large volume regime. The initial state is then obtained by numerically evaluating the following integral
\[
\Psi(v, \phi) = \int dk \Psi(k) e^{-i\alpha} \mathcal{E}_k(v) e^{i\omega(\phi - \phi_0)}
\] (4.10)
with $\Psi(k)$ chosen to be a Gaussian peaked on $k^* = -p^*/\sqrt{12\pi G}$ with a spread $\sigma_k = \sigma/\sqrt{12\pi G}$:
\[
\Psi(k) = e^{-(k-k^*)^2/2\sigma^2}.
\] (4.11)
Further, the time derivative of the initial state can be numerically evaluated as follows
\[
\partial_\phi \Psi(v, \phi) = \int dk i\omega \Psi(k) e^{-i\alpha} \mathcal{E}_k(v) e^{i\omega(\phi - \phi_0)}.
\] (4.12)

### 4.4. Uncertainty products and relative dispersions for method 1, 2 and 3 initial states

The state-dependent properties of the evolution is captured by the product of the uncertainties in $V$ and $p_\phi$ (which are related through the uncertainty relation between $\phi$ and $p_\phi$). An important feature of the choice of initial state is the symmetry of the dispersion of the state across the bounce. It turns out (also noted previously in [3, 4]) that the relative dispersion in $V$ may or may not be symmetric on the two sides of the bounce depending on the type of initial state and the dispersion in $p_\phi$.

In the Wheeler–DeWitt theory the time parameter ‘$\phi$’ is related to the volume $V$ via the following relation\(^{10}\)
\[
\phi = \frac{1}{\sqrt{12\pi G}} \ln \left( \frac{V}{V_0} \right) + \phi_0.
\] (4.13)
Using this relation we can obtain a relation between the dispersion in $\phi$ and the relative dispersion in volume $\Delta V/V$ as follows
\[
\Delta \phi = \frac{1}{\sqrt{12\pi G}} \frac{\Delta V}{V}.
\] (4.14)
The product of the uncertainties of the matter sector is given as
\[
\Delta \phi \Delta p_\phi \geq \frac{\hbar}{2}.
\] (4.15)
Since, the evolution equations are de-parameterized with respect to the scalar field $\phi$ which plays the role of emergent time, we can write the above uncertainty product as the product of the relative dispersion in volume $\Delta V/V$ and the dispersion in the field momentum $\Delta p_\phi$ as follows
\[
\frac{\Delta V}{V} \Delta p_\phi \geq \sqrt{3\pi G}\hbar.
\] (4.16)
Thus, for a minimum uncertainty initial state the above inequality turns to the following equality
\[
\left( \frac{\Delta V}{V} \Delta p_\phi \right)_{\text{min}} = \sqrt{3\pi G}\hbar.
\] (4.17)
\(^{10}\)This result is straightforward to derive using the classical evolution equations. The Wheeler–DeWitt states are peaked on this classical trajectory (see [3, 4, 29] for details).
The initial state for method-2 is a minimum uncertainty state by construction [3], and the above equality holds for all values of $\Delta p \phi$ and $\Delta V/V$. Let us now consider a particular choice $\Delta p \phi$ and $\Delta V/V$ such that the relative dispersions in volume and the field momentum are equal. Together with equation (4.17), this yields the following relation for method-2

$$\frac{\Delta p \phi}{p \phi} = \frac{\Delta V}{V} = \left( \frac{3\pi G \hbar}{p \phi} \right)^{1/2},$$

so that

$$\Delta p \phi = \left( 3\pi p \phi^2 G \hbar^2 \right)^{1/4}, \quad \text{and} \quad \frac{\Delta V}{V} = \left( \frac{3\pi}{p \phi^2 G \hbar^2} \right)^{1/4}. \quad (4.18)$$

It is straightforward to verify that for $\Delta p \phi$ and $\Delta V/V$, the uncertainty product remains minimum during the evolution. For method-1 and method-3 initial data on the other hand, if we choose $\Delta p \phi = \Delta p \phi$ the resulting $\Delta V/V$ will not be equal to $\Delta V/V$, i.e. the state is not a minimal uncertainty state. Further, the numerical simulations show that the following turns out to be true for method-1 and method-3:

- For method-1: the input parameter in this initial data method is $\sigma_v$ which directly determines $\Delta V/V$. Moreover, $\Delta p \phi$ is a non-monotonic function of $\Delta V/V$ and becomes minimum at $\Delta V/V = \left( \frac{3\pi}{p \phi^2 G \hbar^2} \right)^{1/4}$. For method-2, the uncertainty product remains minimum during the evolution. For method-1 and method-3 initial data on the other hand, if we choose $\Delta p \phi = \Delta p \phi$ the resulting $\Delta V/V$ will not be equal to $\Delta V/V$, i.e. the state is not a minimal uncertainty state. Further, the numerical simulations show that the following turns out to be true for method-1 and method-3:

  - For method-1: the input parameter in this initial data method is $\sigma_v$ which directly determines $\Delta V/V$. Moreover, $\Delta p \phi$ is a non-monotonic function of $\Delta V/V$ and becomes minimum at $\Delta V/V = \left( \frac{3\pi}{p \phi^2 G \hbar^2} \right)^{1/4}$.
  - For method-2: the input parameter for this method is $\sigma$ which directly determines $\Delta p \phi$ and $\Delta V/V$ is non-monotonic function of $\Delta p \phi$. It turns out that at $\Delta p \phi = \left( 3\pi p \phi^2 G \hbar^2 \right)^{1/4}$, $\Delta V/V$ is minimized.

The non-monotonic nature of the relative volume dispersion for a Gaussian state was earlier found and studied in [31, 32], where the state considered was different from the initial states discussed in this work. However, the value of $\Delta p \phi$ which minimizes the relative volume dispersion turns out to be the same as we find in our analysis. We will discuss later that $\Delta p \phi$ and $\Delta V/V$ play important roles in understanding the behavior of the deviation of the effective trajectory from the LQC trajectory. It turns out that these values are directly related to the non-monotonic behavior of the quantity that measures the deviation of effective theory from LQC.

5. Implementation of the Chimera scheme

Our goal in this analysis is to consider initial states with a wide variety of values of $p \phi$ and dispersions, including those with small $p \phi$, and those with a large dispersion in volume. Evolution with such states present severe computational challenges. They both require the boundary of the spatial grid to be sufficiently large in order to contain the state within the computational domain during the evolution. Due to the CFL stability condition, for a stable evolution, a larger spatial grid requires a smaller time step [28]. Thus, if the state is widely spread, the time steps in the evolution have to be small, which leads to a larger computation time. Note that the discretization in the quantum difference equation (2.10), denoted by $\Delta v$, is 4 times the Planck volume and is fixed by the underlying quantum geometry. Since the discreteness in the spatial grid cannot be changed, the computational cost grows quadratically with the location of the outer boundary. Therefore, it is difficult to study the numerical evolution of very widely spread states with the numerical approaches used in [2-4]. To circumvent this problem we use a hybrid numerical scheme, named Chimera, which
was introduced by the authors recently [29]. It is an explicit time integration scheme based on the idea of a hybrid spatial grid which takes advantage of the fact that the LQC difference equation can be approximated by the Wheeler–DeWitt equation at large volume. Therefore, we can divide the entire domain into two parts: the inner grid on which we solve the LQC difference equation and an outer grid where we solve the Wheeler–DeWitt equation. On the outer grid we take advantage of the fact that the Wheeler–DeWitt equation has a continuum limit allowing us to transform to numerically better behaved coordinates, so that the time step no longer depends on the outer boundary location. Below we summarize the way this scheme is implemented. A more elaborate discussion of the scheme including several robustness tests has been provided in our previous article, which is focused on the details of the numerical properties of the scheme [29].

Let us denote the boundary of the inner and the outer grids as \( v_{\text{interface}} \). This boundary is chosen at a volume large enough, where the LQC difference equation is extremely well approximated by the Wheeler–DeWitt equation. On the inner grid, where the LQC difference equations are evolved, we use \( v \) as our variable with discreteness \( \Delta v = 4 \). On the outer grid we use \( x = \ln v \) as our variable. This scheme now allows us to place the outer boundary at very large values of \( v_{\text{outer}} = \exp(x_{\text{outer}}) \) without (i) having to use extremely large number of grid points, and (ii) having to use an extremely small time step in \( \phi \). This leads to a very large reduction in computational cost and makes the numerical evolution of states with small \( p_\phi^* \), and those with large dispersion in volume practically feasible.

In the Chimera scheme, it is possible to use different methods to solve the Wheeler—equation, which is a partial differential equation, on the outer grid. In our previous article [29], we introduced two separate implementations of the Chimera scheme: (i) a finite difference (FD) implementation where we use standard centered second order finite differencing on the outer grid, and (ii) a discontinuous Galerkin (DG) implementation where we use the DG method to solve the Wheeler–DeWitt equation on the outer grid. We found that the DG implementation of the Chimera scheme is an order of magnitude more efficient than the FD implementation, since the higher accuracy provided by DG requires less grid points. Using the Chimera scheme we can perform the simulation of a widely spread state with \( p_\phi^* = 20\sqrt{\hbar G} \), which would have taken \( 10^{27} \) years, in 1118 s using the FD implementation with 253, 953 grid points on the outer grid, and in 144 s using the DG implementation with 8125 grid points on the outer grid, while, in both cases, choosing 7500 grid points on the inner grid. A numerical evolution with twice the resolution takes 9148 s using the FD implementation and 472 s using the DG implementation. (For a detailed discussion of the efficiency of the scheme and its performance, see [29]). Since the DG implementation is more efficient than the FD implementation we use it exclusively for the simulations presented here.

6. Evolution of initial states

With the Chimera scheme at our disposal, we can now extend the numerical simulations to initial data parameter regimes, which were not accessible in previous numerical investigations. We have performed a large number of simulations using initial states from the three different initial data methods described in section 4. In this section our goal is to discuss a sample of these results, obtained from evolution of initial states peaked at large and small values of \( p_\phi^* \) and different values of dispersion in volume, as a demonstration of the implementation of the Chimera scheme and to provide a glimpse of some of the results. Detailed discussions of the

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11 In the discussion in this section, \( \Delta \) should not be confused with the symbol used for dispersion elsewhere in this manuscript.
Figure 1. The plot shows the evolution of the expectation value and dispersion of the volume for an initial state peaked at a large value of $p_\phi$: $p_\phi^* = 1500 \sqrt{\hbar G}$ and $\sigma = 95$ for method-3 initial data. The dispersion is depicted with error bars. The bounce happens far away from the Planck volume and the effective trajectory is an excellent approximation to the LQC trajectory.

simulations performed with the different methods and various results is provided in the next section. In the following subsections, we first compare the expectation values of volume in LQC and the effective theory, for the case of initial states constructed with method-3. In the subsequent part of this section, we discuss the results corresponding to some values of $p_\phi^*$ for relative volume dispersion, and the constraints on the fluctuations in pre and post bounce phases for all the methods. As discussed before, the initial state is characterized by three main parameters: the field momentum at which it is peaked $p_\phi^*$, the width of the state in the field momentum $\sigma$ and the volume $v^*$ at which the initial state is peaked. Here we will consider a variety of values of $p_\phi^*$ and $\sigma$ while keeping the value of the volume to be large. A large value of $v^*$ is taken to ensure that the initial state is peaked at a classical trajectory. The value of $p_\phi^*$ and $\sigma$ on the other hand govern whether the state is sharply peaked. The larger the value of $\sigma$, the more sharply peaked the state is in $v^{12}$.

6.1. Expectation value of $V$

Figure 1 shows the evolution of the expectation value and dispersion of the volume observable for the initial state constructed with method-3 peaked at $p_\phi^* = 1500 \sqrt{\hbar G}$ and with $\sigma = 95$. We make a comparison with the trajectory obtained from the effective spacetime description (blue solid curve). It can be seen that the expectation value of the volume is in excellent agreement with the effective evolution throughout the evolution including the bounce point at volume $V_b \approx 1660 l_{Pl}^3$, which according to equation (3.6) occurs at approximately 1658 $l_{Pl}^3$ in the effective theory. In this particular simulation, the properties of the state are compatible with the assumptions made in the derivation of the effective theory—the state is sharply peaked and

$^{12}$ In the following the results are plotted and discussed in terms of the physical volume $V$, whereas for the evolution we use the variable $v$ which is related to $V$ via equation (2.8).
Table 1. The dependence of the value of the bounce volume for different values of $p^*_\phi$ on the spread $\sigma$. The bounce volume $V_b$ in the quantum theory is in the Planck units.

| $p^*_\phi = 20\sqrt{G}\hbar$ | $p^*_\phi = 80\sqrt{G}\hbar$ | $p^*_\phi = 140\sqrt{G}\hbar$ | $p^*_\phi = 200\sqrt{G}\hbar$ |
|-----------------------------|-----------------------------|-----------------------------|-----------------------------|
| $\sigma$ | $V_b$ | $\sigma$ | $V_b$ | $\sigma$ | $V_b$ | $\sigma$ | $V_b$ |
| 2.25 | 146.22 | 5.0 | 129.07 | 10.0 | 170.12 | 16.0 | 229.41 |
| 2.5 | 102.66 | 6.0 | 115.03 | 13.0 | 163.69 | 19.0 | 226.97 |
| 2.75 | 79.03 | 7.0 | 107.31 | 15.0 | 161.43 | 22.0 | 225.47 |
| 3.9 | 42.25 | 9.0 | 99.46 | 17.0 | 159.94 | 26.0 | 224.22 |

the volume never gets close to the Planck volume during evolution. Therefore, the agreement of the LQC evolution with the effective dynamical trajectory is expected.

We now discuss the cases when we decrease the value of $p^*_\phi$. We choose two representative values $p^*_\phi = 200\sqrt{G}\hbar$ and $p^*_\phi = 20\sqrt{G}\hbar$. In these simulations, the spread $\sigma$ for each of these values is kept in the range of 8%–15% of $p^*_\phi$. We find that a smaller value of $p^*_\phi$ leads to a smaller bounce volume, and as the spread in $p^*_\phi$ decreases, the relative spread in volume of the state increases, leading to deviation from sharp peakedness. Figure 2 (corresponding to $p^*_\phi = 200\sqrt{G}\hbar$) and figure 3 (corresponding to $p^*_\phi = 20\sqrt{G}\hbar$) show the evolution of the expectation value and dispersion of the volume observable, and the respective effective trajectories. For these particular cases, as the value of $p^*_\phi$ is decreased, the deviation between the effective and LQC trajectory increases. We will later see that this is not a generic feature independent of the initial state construction method. The differences between the effective and the LQC trajectory are in particular quite prominent for $p^*_\phi = 20\sqrt{G}\hbar$, where the value of the spread $\sigma$ in our simulations ranges between 2.25 and 3.0. The volume at the bounce for this $p^*_\phi$, in the effective theory, is $V_{b}^{(\text{eff})} \approx 22 l_P^3$, whereas the bounce volume for LQC trajectories are larger. We make two important observations based on these simulations: (i) there are dispersion dependent deviations in the LQC trajectory from the effective trajectory, and (ii) the deviations seem to grow as the spread $\sigma$ is decreased for a fixed $p^*_\phi$. In the next section, we will discuss the way these results are affected with different choices of initial state construction. For a quick comparison see table 2. Note that the LQC trajectories shown in figures 2 and 3, correspond to a fixed $p^*_\phi$ with different values of $\Delta p_\phi$ such that $\Delta p_\phi < \tilde{\Delta} p_\phi$, defined via equation (4.19). In this regime, as the value of $\sigma$ decreases, $\Delta p_\phi$ decreases which in turn causes the relative volume dispersion to increase and leads to a larger deviation from the effective trajectory.

In table 1, we provide the values of the bounce volume for different values of the field momentum $p^*_\phi$ and spread $\sigma$. It is clear that the bounce volume depends more strongly on the spread of the state for small values of $p^*_\phi$.

6.2. Relative dispersion $\Delta \phi$

In this subsection we study the relation between the relative volume dispersion and the relative dispersion in the field momentum, on the two sides of the bounce. We will again consider the evolution of states with a wide variety of initial conditions. These give rise to a wide range of initial values of the relative dispersion. The study of the evolution of the dispersion of the states are important for several reasons, the most important one being the issue of semi-classicality in the low curvature regime on the two sides of the bounce [30, 31, 43]. These issues can also be analytically understood using triangle inequalities derived in [30]. A comparison of our results with the constraints from triangle inequalities are discussed in the next subsection.
Figure 2. The expectation value of the volume $V$ and the associated dispersion $\Delta V$ (shown as errorbars), for $p_\phi^* = 200\sqrt{\hbar G}$ for different values of $\sigma$ is plotted for method-3 initial data in panel (a). We find that the bounce happens much closer to the Planck volume than in figure 1 (for $p_\phi^* = 1500\sqrt{\hbar G}$) and the effective trajectory shows small deviations from the LQC trajectory (larger for smaller values of $\sigma$). In panel (b), we see (for $\sigma = 16$ only) that although there are differences between the LQC and effective trajectories, the effective theory lies within the dispersions. These trajectories correspond to $\Delta p_\phi < \Delta p_\phi$, for which $\sigma < \widetilde{\sigma} = \sqrt{2}\Delta p_\phi = 35.04$ for $p_\phi^* = 200\sqrt{\hbar G}$. In this regime, increasing $\Delta V/V$ (decreasing $\sigma$) results in larger deviation from the effective theory, as discussed further in table 2.

Let us first discuss the evolution of $\Delta V/V$ in LQC. Figure 4(a) shows the evolution of the relative volume dispersion for a state which is initially sharply peaked on a classical trajectory. The value of the field momentum is $p_\phi^* = 1500\sqrt{\hbar G}$ for all initial data methods and $\sigma_v = 1000$ for method-1 while $\sigma = 95$ for method-2 and 3. From the figure it is clear that far from bounce in the low curvature regime, $\Delta V/V$ tends to a constant value in the asymptotic limit. In
Figure 3. The expectation value of the volume $V$ and the associated dispersion $\Delta V$ (shown as errorbars), for $p_\phi^* = 20\sqrt{G^*}\hbar$ for different values of $\sigma$ is plotted for method-3 initial data in panel (a). We find that the bounce happens much closer to the Planck volume than in figure 1 (for $p_\phi^* = 1500\sqrt{G^*}\hbar$) and there are significant differences between the effective trajectory and the LQC trajectories for various field dispersions (larger for smaller values of $\sigma$). Panel (b) shows the plot of the expectation value of volume, associated volume dispersion, and the corresponding effective trajectory for $\sigma = 2.25$. It is evident that although there are differences between these trajectories, the effective theory lies within $\Delta V$. These trajectories correspond to $\Delta p_\phi < \bar{\Delta}p_\phi$, for which $\sigma < \tilde{\sigma} = \sqrt{2}\bar{\Delta}p_\phi = 11.08$ for $p_\phi^* = 20\sqrt{G^*}\hbar$. As discussed in table 2, increasing $\Delta V/V$ (decreasing $\sigma$) results in larger deviation from the effective theory, in this regime.
method-2 and method-3, as the state is evolved backwards, and the Planckian curvature is reached, the relative volume dispersion decreases. As the evolution is continued backwards, it achieves a minimum value at some point and then starts increasing again. For method-1 initial data, $\Delta V/V$ monotonically increases when evolved backwards. On the other side of the bounce, as the low curvature regime is reached once again, the relative volume dispersion tends to a constant value. The asymptotic constant values on the two sides of the bounce, are equal if the initial state is chosen according to method-3. For method-1 and method-2 initial data, they are different, in general. However, it turns out that for method-2, the asymptotic values

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure4}
\caption{Evolution of the relative dispersion in volume $\Delta V/V$. Panel (a) shows the evolution for $p_\phi = 1500 \sqrt{G\hbar}$ for initial data method-1 ($\sigma_v = 1000$), method-2 and method-3 (both with $\sigma = 95$). Panel (b) shows the evolution of $\Delta V/V$ for $p_\phi = 20 \sqrt{G\hbar} \sigma = 2.5$ for initial data method-2 and 3.}
\end{figure}
of the relative volume dispersion can be brought closer to each other by choosing appropriate values of the relative dispersion in the field momentum.

Figure 4(b) shows the evolution of the relative volume dispersion \( \frac{\Delta V}{V} \) for \( p_0^* = 20\sqrt{\hbar}G \) and \( \sigma = 2.5 \) for method-2 and 3\(^{13} \). As can be seen from the figure, the initial value of the fractional dispersion in volume is as large as 4.3. Due to such a large dispersion in volume, the state is not sharply peaked. However, the evolution across the bounce shows that the relative volume dispersion again takes a constant value in the asymptotic limit on both sides of the bounce. It is important to note that in both cases, i.e. as shown in figures 4(a) and (b), \( \frac{\Delta V}{V} \) approaches a constant value in the asymptotic limit, far from the bounce. Also, the value of the relative dispersion near the bounce is smaller than these asymptotic values for method-2 and 3 initial data.

### 6.3. Triangle inequalities of the dispersion

Let us now turn to the discussion of constraints on the difference of the relative volume dispersion on the two sides of the bounce. A large disparity between the dispersion on the two sides may imply the loss of semi-classicality as the universe evolves from one side of the bounce to the other. Analytical investigations in this direction \([30, 31, 43]\), however, establish that the state of the universe remains semi-classical if one starts with a semi-classical state on one side of the bounce and evolves it across. According to \([30]\) the asymptotic values of the relative dispersions, far from the bounce obey triangle inequalities given as follows

\[
\Sigma_- \leq \Sigma_+ + 2 \Sigma \quad \text{and} \quad \Sigma_+ \leq \Sigma_- + 2 \Sigma, \quad (6.1)
\]

where \( \Sigma_+ \) and \( \Sigma_- \) are the asymptotic values of the relative volume dispersion in the expanding and contracting branches respectively and \( \Sigma = \Delta p_0/p_0 \) is the relative dispersion in the field momentum. The triangle inequalities given in equation (6.1), in a sense, place bounds on the asymmetry in the relative volume dispersion on the two sides of the bounce

\[
|\Sigma_- - \Sigma_+| \leq 2\Sigma. \quad (6.2)
\]

Figures 5–7 show the evolution of the relative volume dispersion for method-1, 2 and 3 respectively, with large and small values of the field momentum. For method-2 and 3 we show the simulations for \( p_0^* \) as small as \( 20\sqrt{\hbar}G \), whereas for method-1 we restrict ourselves to \( p_0^* \geq 500 \) (see footnote 13). The dotted and dashed curves show \( \Sigma_\pm + 2 \Sigma \). The figures clearly show that on both sides of the bounce, the triangle inequalities are satisfied i.e. \( \Sigma_- < \Sigma_+ + 2 \Sigma \). It is also important to note that these inequalities holds true for all three initial data methods. Figure 8 shows the asymptotic values of the volume dispersion, where \( \Sigma_\pm \) is compared with \( \Sigma_\mp + 2 \Sigma \), for method-1 initial data. Similarly, figures 9 and 10 show the asymptotic values of the relative volume dispersion, where \( \Sigma_\pm \) is compared with \( \Sigma_\mp + 2 \Sigma \), for method-2 and method-3 initial data. The dashed curves in the upper panel of these figures show \( \Sigma_- \) and in the lower panel \( \Sigma_+ \) for various values of \( p_0^* \).

### 7. Results

In this section, we present results based on a large number of simulations by considering states peaked at various values of \( p_0^* \) in the range \( 20\sqrt{\hbar}G < p_0^* < 1500\sqrt{\hbar}G \) for method-2 and 3 initial data, and for \( p_0^* \geq 500\sqrt{\hbar}G \) for method-1. For each value of \( p_0^* \) we vary \( \Delta p_0 \) (the

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\(^{13}\) For method-1, there are difficulties associated with a reliable construction of initial states for small values of \( p_0^* \). These difficulties seem to be independent of the numerical procedure used for the evolution. Due to this reason, in our analysis small values of field momentum are not discussed for method-1, and we restrict ourselves to \( p_0^* \geq 500 \).
Figure 5. Triangle inequalities for method-1 initial data for $p^*_\phi = 1500\sqrt{\frac{G}{\hbar}}$ (panel (a)) and $p^*_\phi = 500\sqrt{\frac{G}{\hbar}}$ (panel (b)) and $\sigma_v = 1000$ are shown to be valid throughout the evolution.

spread of the Gaussian) in the range 5% to 15% of the mean value of the field momentum $p^*_\phi$. The value of $p^*_\phi$ determines the bounce volume while $\Delta p_\phi$ dictates whether the state is sharply peaked initially. For each of the simulations we compute the expectation values of various quantities and their relative dispersions both at the bounce and in the asymptotic limits far from the bounce. This way we are able to make a detailed quantitative investigation of the spread dependent properties of the initial state during evolution. The analysis performed here enables us to explore in a rigorous way the deviation of the LQC trajectory from the effective
Figure 6. This figure shows the evolution of the relative volume dispersion and the validity of the triangle inequality for $p^*_\phi = 1500\sqrt{G\hbar}$ (panel (a)) and $p^*_\phi = 20\sqrt{G\hbar}$ (panel (b)) and $\sigma = 95$ for method-2 initial data. We find that the triangle inequalities are valid throughout the evolution for both large and small $p^*_\phi$.

trajectory in a quantitative fashion. The study of the spread dependent corrections provides us valuable insights into which regime the effective theory is reliable.

As in equation (3.7), in order to quantify the deviation of LQC trajectory from the effective dynamical trajectory, we define the quantity

$$\delta = \frac{V_b - V_b^{(\text{eff})}}{V_b^{(\text{eff})}},$$

where $V_b$ denotes the expectation value of the bounce volume in the LQC trajectory and $V_b^{(\text{eff})}$ is the bounce volume in the effective trajectory calculated using equation (3.6) by setting $p_b = p^*_\phi$ (the field momentum at which the initial state is peaked). The quantity $\delta$ denotes the
fractional difference in the bounce volumes in LQC and effective theory with respect to the bounce volume in the effective theory.

Before we discuss the results from each of the methods, we summarize the key results regarding the dependence of the quantity \( \delta \) on the initial data parameters in table 2. It is evident from the table that \( \delta \) depends on several different factors including \( p_\phi^* \), \( \Delta p_\phi \) and \( \Delta V/V \) and the initial data method. For example, if one fixes the relative volume dispersion and increases the field momentum \( p_\phi^* \), then \( \delta \) decreases for method-1 and method-2 for all \( \Delta p_\phi \), whereas...
Figure 8. The validity of the triangle inequalities for method-1 initial data for varying $\Delta p_\phi$: panel (a) and (b) show the variation of $\Sigma_+ + 2\Sigma$ and $\Sigma_- + 2\Sigma$, respectively, demonstrating the validity of the first and the second inequality in equation (6.1). The dashed curves in the upper panel show the value of $\Sigma_-$ for different values of $p_\phi^*$. The upper most dashed curve correspond to $p_\phi^* = 500 \sqrt{G\hbar}$ and as $p_\phi^*$ increases the corresponding dashed curve for $\Sigma_-$ shifts downwards. Similarly, in the lower panel, the dashed curves correspond to $\Sigma_+$ for different values of $p_\phi^*$. The left most curve corresponds to $p_\phi^* = 500 \sqrt{G\hbar}$ and the right most to $p_\phi^* = 1500 \sqrt{G\hbar}$.

for method-3, $\delta$ increases if $\Delta p_\phi < \Delta \tilde{p}_\phi$. Similarly, for a fixed value of the field momentum, $\delta$ decreases with increasing $\Delta V/V$ for all $\Delta p_\phi$ for method-1, whereas for method-2 and 3, the behavior is different for the two regimes $\Delta p_\phi < \Delta \tilde{p}_\phi$ and $\Delta p_\phi > \Delta \tilde{p}_\phi$. Note that, for method-1, the parameters of the initial data are different compared to method-2 and 3. The initial data in method-1 is a Gaussian which is characterized by $\sigma_v (= \sqrt{2}\Delta V)$, $v_*$ and $p_\phi^*$, and the value of $\Delta p_\phi$ is then derived from these. On the other hand, for method-2 and
Figure 9. This figure shows the validity of the triangle inequalities for initial data of type method-2 for varying $\Delta p_\phi$. Panel (a) and (b) show the variation of $\Sigma_+ + 2\Sigma$ and $\Sigma_- + 2\Sigma$, respectively, demonstrating the validity of the first and the second inequality in equation (6.1). It is interesting to note that the inequalities are saturated in the low $\Delta p_\phi$ region.

Table 2. Summary of the main results.

| Initial data | Fixed $\frac{\Delta \Sigma}{\Sigma}$ increasing $p_\phi^*$ | Fixed $p_\phi^*$ increasing $\frac{\Delta \Sigma}{\Sigma}$ | Fixed $\Delta p_\phi$ increasing $p_\phi^*$ |
|--------------|----------------------------------------------------------|--------------------------------------------------------|-------------------------------------------|
| Method-1     | $\delta$ decreases                                      | $\delta$ decreases                                      | $\delta$ decreases if $\frac{\Delta \Sigma}{\Sigma} < \frac{\Delta p_\phi}{p_\phi}$ |
|              |                                                          | $\delta$ increases if $\frac{\Delta \Sigma}{\Sigma} > \frac{\Delta p_\phi}{p_\phi}$              | $\delta$ decreases if $\frac{\Delta \Sigma}{\Sigma} > \frac{\Delta p_\phi}{p_\phi}$ |
| Method-2     | $\delta$ decreases                                      | $\delta$ increases if $\Delta p_\phi < \Delta p_\phi^*$ | $\delta$ decreases if $\Delta p_\phi > \Delta p_\phi^*$ |
|              | $\delta$ increases if $\Delta p_\phi < \Delta p_\phi^*$ |                                                          | $\delta$ decreases if $\Delta p_\phi > \Delta p_\phi^*$ |
| Method-3     | $\delta$ increases if $\Delta p_\phi < \Delta p_\phi^*$ | $\delta$ increases if $\Delta p_\phi < \Delta p_\phi^*$ | $\delta$ decreases if $\Delta p_\phi > \Delta p_\phi^*$ |
|              | $\delta$ decreases if $\Delta p_\phi > \Delta p_\phi^*$ |                                                          | $\delta$ decreases if $\Delta p_\phi > \Delta p_\phi^*$ |
|              | $\delta$ is independent of $p_\phi$                     |                                                        | $\delta$ is independent of $p_\phi$ |
method-3 the parameters of the initial data are $\sigma (= \sqrt{2} \Delta p_\phi)$, $v_*$ and $p_0^*$. In the following, we consider the results for the three different initial data methods in separate subsections.

7.1. Method-1: Gaussian state

(1) As discussed previously in section 4, for method-1 initial data, $\Delta V/V$ appears as an input parameter and the value of the dispersion in the field momentum is computed from it. It turns out that $\Delta p_\phi$ depends non-monotonically on $\Delta V/V$ as can be seen from figure 11(a) which shows the initial field dispersion $\Delta p_\phi$ plotted against the initial relative volume dispersion far from the bounce in the expanding branch. The turnaround in
Figure 11. Panel (a) shows the behavior of $\Delta p_\phi$ plotted against $\Delta V/V$ for method-1 initial data. It is clear that $\Delta p_\phi$ shows non-monotonic behavior with $\Delta V/V$. In panel (b), $\Delta V/V \sqrt{p_\phi}$ is plotted against $\Delta p_\phi$, in order to make it clear that irrespective of the value of $p_\phi$, the minimum of $\Delta p_\phi$ occurs when $\frac{\Delta V}{V} \sqrt{p_\phi} = (3\pi G \hbar)^{1/4}$.

Due to this non-monotonic dependence of $\Delta p_\phi$ on $\Delta V/V$ any quantity varying monotonically with respect to $\Delta V/V$ will vary non-monotonically with respect to $\Delta p_\phi$.

Figure 12 shows the variation of $\delta$ for different values of $p_\phi$ as function of the initial relative dispersion in volume $\Delta V/V$ for method-1 initial data. As $\Delta V/V$ increases for a fixed value of $p_\phi$, $\delta$ decreases monotonically. This behavior is summarized in the row for

\[ p_\phi = 500, 750, 1000, 1250, 1500 \]
Figure 12. The variation of the quantity $\delta$ is plotted against the relative dispersion in volume $\Delta V/V$ for method-1 initial data. For all data points, the state is initially peaked at $v^\ast = 20000$. The different lines correspond to different values of the field momentum $p_\phi^\ast$ in units of $\sqrt{G\hbar}$.

Figure 13. This plot shows the behavior of the quantity $\delta$ with respect to the field momentum $\Delta p_\phi$ for method-1 initial data for several different values of $p_\phi^\ast$. The value of $v^\ast$ is the same as in figure 12.

method-1 and the column with fixed $p_\phi^\ast$ in table 2. This presents a counter example to the expectation that a wide-spread state should show larger differences between the effective and the LQC trajectory than a narrow state. It is also evident from the figure that for a fixed value of $\Delta V/V$, a lower value of $p_\phi^\ast$ results in a larger value of $\delta$ (summarized in the row for method-1 and column for fixed $\Delta V/V$ in table 2). The variation of $\delta$ with respect to $\Delta p_\phi$ is shown in figure 13. Here we see a clear non-monotonic behavior. The
(3) As discussed before, the scalar field $\phi$ plays the role of internal time in our analysis. Its absolute value does not have a physical meaning as it can be shifted without affecting the physics. However, a ‘time’ difference is a useful quantity to study. We consider here the difference between the values of the scalar field at the bounce in the effective and the LQC trajectory, i.e. $(\phi_b - \phi_b^{(eff)})$, and plot it against $\Delta p_\phi$ in figure 14. It is evident that the LQC trajectory bounces earlier than the effective one and the difference in bounce time increases with increasing $\Delta p_\phi$ if $\Delta V/V < \bar{\Delta V}/V$ and decreases with increasing $\Delta p_\phi$ if $\Delta V/V > \bar{\Delta V}/V$. Moreover, for a fixed $\Delta p_\phi$ this difference decreases with increasing $p_\phi^*$ if $\Delta V/V < \bar{\Delta V}/V$ and increases with $p_\phi^*$ if $\Delta V/V > \bar{\Delta V}/V$.

(4) The energy density at the bounce $\rho_b$ also depends on the relative volume dispersion of the initial state $\Delta V/V$. As shown in figure 15(a), for a fixed value of $p_\phi^*$, $\rho_b$ decreases with a decreasing spread in the volume. On the other hand, if the dispersion in volume is fixed, then $\rho_b$ takes smaller value as $p_\phi^*$ decreases. It is to be noted that the energy density never exceeds the upper bound on $\rho_b$, i.e. $\rho_b \leq \rho_{\text{max}} \approx 0.409 \rho_0$ derived from the exactly solvable model in [19] and which is also predicted by the effective theory equation (3.4). The deviation of $\rho_b$ from $\rho_{\text{max}}$ is also a measure of the deviation of the effective theory from LQC. That is, the larger the difference between $\rho_b$ and $\rho_{\text{max}}$, the larger is the departure between the LQC and effective trajectory and the larger is $\delta$. Figure 15(b) shows the variation of $\rho_b$ with respect to the dispersion in the field momentum $\Delta p_\phi$. The inset in this figure shows the non-monotonic variation of $\rho_b$ with $\Delta p_\phi$. It can also be seen that the energy density always remains below the critical value for all values of the field momentum.
7.2. Method-2: Wheeler–DeWitt initial state

Figure 16 shows the variation of the initial relative volume dispersion $\Delta V/V$ with varying initial spread in the field momentum $\Delta p_\Phi$, far from bounce in the expanding branch. For the entire range of $\Delta p_\Phi$, $\Delta V/V$ decreases monotonically with increasing $\Delta p_\Phi$. This implies that if a quantity varies monotonically with respect to $\Delta p_\Phi$, it should do so with...
Figure 16. Plot of the variation in $\Delta V/V$ with respect to $\Delta p_\phi$ for method-2 initial data for different values of $p_\phi^*$. It is clear that the relative volume dispersion varies with $\Delta p_\phi$ in a monotonic fashion. The values of the field momentum are given in the units of $\sqrt{G\hbar}$.

Figure 17. The figure shows the variation of $\delta$ with respect to the spread in the field momentum $\Delta p_\phi$ for method-2 initial data, in a log–log plot. It is evident from the figure that in the low $\Delta p_\phi$ regime, $\delta$ is large, and as $\Delta p_\phi$ is increased, $\delta$ initially decreases monotonically. Then, around $\Delta p_\phi \approx 10$, there is a small interval of almost linear behavior. As $\Delta p_\phi$ is increased further, the curves for

respect to $\Delta V/V$ as well. Note that a method-2 initial state is a minimal uncertainty state in $(\phi, p_\phi)$, and the variation of relative volume dispersion with respect to the dispersion in field momentum is monotonic. This is not the case for method-1 and 3 initial state, which are not minimal uncertainty states in $(\phi, p_\phi)$.

(2) Figure 17 shows the variation of $\delta$ with respect to $\Delta p_\phi$ for method-2 initial data, in a log–log plot. It is evident from the figure that in the low $\Delta p_\phi$ regime, $\delta$ is large, and as $\Delta p_\phi$ is increased, $\delta$ initially decreases monotonically. Then, around $\Delta p_\phi \approx 10$, there is a small interval of almost linear behavior. As $\Delta p_\phi$ is increased further, the curves for
different values of \( p^*_\phi \) start to deviate from each other, reach a minimum value (at a \( p^*_\phi \) dependent value of \( \Delta p_\phi \)) and then start to increase again. In this way, the curve for \( \delta \) shows non-monotonicity with respect to the dispersion in the field momentum in the large \( \Delta p_\phi \) regime. It is noteworthy that the minimum of \( \delta \) occurs at \( \Delta p_\phi = \Delta \tilde{p}_\phi = (3\pi p^*_\phi^2 G \hbar^2)^{1/4} \). This figure also shows that for a fixed value of \( \Delta p_\phi \), the value of \( \delta \) decreases with increasing \( p^*_\phi \). This behavior is summarized in the row for method-2 and column for fixed \( \Delta p_\phi \) in table 2.

(3) Figure 18(a) shows the behavior of \( \delta \) with varying relative dispersion in volume \( \Delta V/V \) for different values of \( p^*_\phi \) and \( \Delta p_\phi \). Each curve corresponds to different values of \( p^*_\phi \) with the data points corresponding to different \( \Delta p_\phi \). We find that in the large \( \Delta V/V \) regime \( \delta \) is large. As the value of \( \Delta V/V \) is decreased, \( \delta \) decreases until it reaches a minimum value at \( \Delta V/V = (3\pi G \hbar^2 / p^*_\phi)^{1/4} \) after which it starts to increase again. Figure 18(b) shows the behavior of \( \delta \) zoomed in near the location of the minima to highlight the non-monotonicity. The dashed curve joining the separate \( p^*_\phi \) curves correspond to the points where \( \delta \) is minimum and \( \Delta p_\phi = \Delta \tilde{p}_\phi = (3\pi p^*_\phi^2 G \hbar^2)^{1/4} \). To the left of the dashed curve \( \Delta p_\phi > \Delta \tilde{p}_\phi \) and \( \delta \) decreases with increasing \( \Delta V/V \), while to the right of the dashed curve \( \Delta p_\phi < \Delta \tilde{p}_\phi \) and \( \delta \) increases with increasing \( \Delta V/V \). This behavior is summarized in the row for method-2 and the column for fixed \( p^*_\phi \) in table 2. It can also be seen that \( \delta \) decreases with increasing \( p^*_\phi \) for a fixed value of \( \Delta V/V \) as summarized in the row for method-2 and the column for fixed \( \Delta V/V \) in table 2.

(4) Figure 19 shows the variation of the difference in the bounce time between the LQC and effective trajectories \( (\phi_b - \phi_b^{(\text{eff})}) \) with varying \( \Delta p_\phi \) in a log–log plot. It is clear that in the low \( \Delta p_\phi \) regime, the curve is linear, which implies a power-law relation between \( (\phi_b - \phi_b^{(\text{eff})}) \) and \( \Delta p_\phi \)

\[
(\phi_b - \phi_b^{(\text{eff})}) \propto (\Delta p_\phi)^\nu, \tag{7.1}
\]

where \( \nu \) is an exponent. A numerical fit to the curve in the linear region gives the value of the exponent to be \( \nu \approx -2.017 \pm 0.008 \). As we will see in the next subsection, this power law also holds true for method-3 initial data, with the same exponent to three significant figures. In the large \( \Delta p_\phi \) region there is non-monotonic behavior and \( (\phi_b - \phi_b^{(\text{eff})}) \) reaches a minimum at \( \Delta p_\phi = \Delta \tilde{p}_\phi = (3\pi p^*_\phi^2 G \hbar^2)^{1/4} \).

(5) Figure 20(a) and (b) show the variation in the energy density at the bounce with respect to the relative volume dispersion \( \Delta V/V \) and the field dispersion \( \Delta p_\phi \), respectively. It is evident from the figure, that the energy density at the bounce is very close to the maximum allowed energy density \( \rho_{\text{max}} = 0.409 \rho_{\text{Pl}} \) for sharply peaked states (corresponding to small \( \Delta V/V \)). It is, however, noteworthy that the energy density at the bounce always remains smaller than \( \rho_{\text{max}} \). A similar dependence of the energy density at the bounce is discussed in [31] when the wavefunction is taken to be a squeezed state. There, it was found that increasing squeezing leads to an increasing \( \Delta V/V \), which decreases \( \rho_b \).

### 7.3. Method-3: rotated Wheeler–DeWitt initial state

(1) Figure 21(a) shows the behavior of the relative dispersion in volume \( \Delta V/V \) with respect to \( \Delta p_\phi \) in the regime far from the bounce in the expanding branch. It is evident that in the large \( \Delta p_\phi \) regime, the relative volume dispersion has a minimum at a certain value of \( \Delta p_\phi \). This minimum occurs at \( \Delta p_\phi = \Delta \tilde{p}_\phi = (3\pi p^*_\phi^2 G \hbar^2)^{1/4} \). As discussed in section 4 this is exactly the location at which \( \Delta V/V \) as a function of \( \Delta p_\phi \) has a minimum for method-3 initial data. This is clarified in figure 21(b) that shows \( \Delta p_\phi \) plotted against \( \Delta V/V \).
Figure 18. The quantity $\delta$ plotted with respect to the relative dispersion in volume $\Delta V/V$ for method-2 initial data. It is clear from figure 18(a) that $\delta$ increases with increasing $\Delta V/V$ in the large $\Delta V/V$ regime. Figure 18(b) shows the same plot zoomed into the small $\Delta V/V$ regime, where the non-monotonicity of $\delta$ is apparent. The dashed curve in figure 18(b) intersects the curves with various values of $p_0^\phi$ at the points where the relative fluctuation in $p_\phi$ and $V$ are equal, i.e. $\frac{\Delta p_\phi}{p_\phi} = \frac{\Delta V}{V} = \left(\frac{2\pi m}{\hbar^2}\right)^{1/2}$ (see equation (4.18)).

(2) Figure 22 shows the variation of $\delta$ with respect to $\Delta p_\phi$ for several values of $p_0^\phi$. It is interesting to note that $\delta$ varies monotonically with $\Delta p_\phi$. As can be seen, the curve is linear in the large $\Delta p_\phi$ regime, where the initial state is relatively sharply peaked. This indicates a power law relation between $\delta$ and $\Delta p_\phi$

$$\delta \propto (\Delta p_\phi)^\mu,$$  \hspace{1cm} (7.2)
where $\mu$ is a constant exponent. A numerical fit yields the value to be $\mu = -2.018 \pm 0.001$ (in the large $p_0^*$ regime). As we will discuss later in this subsection, a similar power-law relation is also present in the behavior of the difference in bounce time between the LQC and effective trajectories as a function of $\Delta p_\phi$. In the small $\Delta p_\phi$ region the curve tilts slightly upward compared to straight line behavior.

(3) Figure 23(a) shows the variation of $\delta$ with respect to the relative dispersion in volume $\Delta V/V$. The value of $\delta$ behaves almost linearly in the small $p_0^*$ regime where $\Delta V/V$ is large. As the relative dispersion in $V$ decreases, i.e. in the large $p_0^*$ regime, and the state becomes more sharply peaked, the quantity $\delta$ decreases indicating that the effective theory serves as a better approximation to the full LQC evolution in this case with decreasing $\Delta V/V$. Figure 23(b) shows the behavior of $\delta$ with the relative volume dispersion $\Delta V/V$, in the regime corresponding to small $\Delta V/V$ and large $p_0^*$. Different curves in figure 23(b) show the variation of $\delta$ for different $p_\phi^*$ and the data points on each curve represent different $\Delta p_\phi$. Since $\Delta V/V$ varies non-monotonically with $\Delta p_\phi$ in this regime, the behavior of $\delta$ with varying $\Delta V/V$ is also non-monotonic. Recall that the input parameter for method-3 initial data are $p_\phi^*$ and $\Delta p_\phi$, hence $\Delta V/V$ is a derived quantity. The dashed curve connecting the turnaround points of different curves corresponds to $\Delta p_\phi = \Delta \tilde{p}_\phi = (3\pi^2 G\hbar^2)^{1/4}$ which is where $\Delta V/V$ is minimum and $\delta$ changes behavior. Below this curve ($\Delta p_\phi > \Delta \tilde{p}_\phi$) $\delta$ decreases with increasing $\Delta V/V$, while above the curve ($\Delta p_\phi < \Delta \tilde{p}_\phi$) $\delta$ increases. This is summarized in the row for method-3 and the column for fixed $p_\phi^*$ in table 2. The figure also shows that for a fixed value of $\Delta V/V$, the quantity $\delta$ increases with increasing $p_\phi^*$ above the dashed curve ($\Delta p_\phi < \Delta \tilde{p}_\phi$) while $\delta$ decreases with increasing $p_\phi^*$ below the dashed curve ($\Delta p_\phi > \Delta \tilde{p}_\phi$). This is summarized in the row for method-3 and the column for fixed $\Delta V/V$ in table 2. Figure 24 shows the dependence of $\delta$ with varying $p_\phi^*$ for different values of $\Delta p_\phi$. It can be seen that $\delta$ is practically independent of $p_\phi^*$ for a given

![Figure 19](image_url)

Figure 19. The variation of the difference in the time of bounce in the LQC evolution and the effective dynamical evolution as a function of $\Delta p_\phi$ for different values of $p_\phi^*$ for method-2 initial data is shown. Clearly, in the small $\Delta p_\phi$ regime, the curve is linear, implying a power law relation between $(\phi_h - \phi_e)$ and $\Delta p_\phi$ given in equation (7.1). The slope of the curves is $\nu = -2.017 \pm 0.008$. The variation of the difference in the time of bounce in the LQC evolution and the effective dynamical evolution as a function of $\Delta p_\phi$ for different values of $p_\phi^*$ for method-2 initial data is shown. Clearly, in the small $\Delta p_\phi$ regime, the curve is linear, implying a power law relation between $(\phi_h - \phi_e)$ and $\Delta p_\phi$ given in equation (7.1). The slope of the curves is $\nu = -2.017 \pm 0.008$. The variation of the difference in the time of bounce in the LQC evolution and the effective dynamical evolution as a function of $\Delta p_\phi$ for different values of $p_\phi^*$ for method-2 initial data is shown. Clearly, in the small $\Delta p_\phi$ regime, the curve is linear, implying a power law relation between $(\phi_h - \phi_e)$ and $\Delta p_\phi$ given in equation (7.1). The slope of the curves is $\nu = -2.017 \pm 0.008$.
value of $\Delta p_\phi$ as summarized in the row for method-3 and the column for fixed $\Delta p_\phi$ in table 2.

(4) Figure 25 shows the variation of the difference in the values of emergent time ‘$\phi_b$’ at the bounce in the LQC and effective trajectories ($\phi_b - \phi_b^{(\text{eff})}$) with respect to $\Delta p_\phi$. There is a power law relation between these quantities over the complete range of values of $\Delta p_\phi$ considered:

$$(\phi_b - \phi_b^{(\text{eff})}) \propto (\Delta p_\phi)^\nu,$$

(7.3)

with $\nu$ being the power law exponent, which turns out to be $-2.015 \pm 0.003$. This power law is very similar to the power law relating $\delta$ to $\Delta p_\phi$ as given in equation (7.2) and even
Figure 21. Panel (a) shows the variation in the value of $\Delta V/V$ with $\Delta p_\phi$ for method-3 initial data in the regime far from the bounce in the expanding branch. It is evident that $\Delta V/V$ varies non-monotonically with increasing $\Delta p_\phi$. Panel (b) shows $\Delta p_\phi/\sqrt{p_\phi}$ plotted against $\Delta V/V$. We find that $\Delta p_\phi/\sqrt{p_\phi} = (3\pi G \hbar^2)^{1/4}$ when $\Delta V/V$ is minimal, as discussed in section 4.

has the same exponent $\mu = \nu = -2.01$ to three significant figures. Also, $\nu$ is surprisingly the same (within the fitting errorbars) as the exponent in the power law behavior of $(\phi_b - \phi_b^{(eff)})$ for method-2, which is $-2.017 \pm 0.008$.

(5) Figure 26 shows the variation of the energy density at the bounce $\rho_b$ with respect to $\Delta V/V$ (upper panel) and $\Delta p_\phi$ (lower panel). As the relative dispersion in volume increases, the energy density at the bounce decreases. In the small $\Delta V/V$ regime, where the state is sharply peaked in volume, the energy density is close to but smaller than $\rho_{\text{max}}$. 

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8. Discussion

In this paper we have presented a rigorous numerical study of the evolution of states in LQC for a flat FRW universe in the presence of a massless scalar field, including states which are widely spread and bounce close to the Planck volume. Due to the underlying discrete quantum geometry, the evolution of states is non-singular and the big bang singularity is replaced by a quantum bounce. Previous numerical studies in LQC demonstrated the existence of a bounce for sharply peaked states corresponding to large values of the field momentum $p_\phi$. Such states lead to a bounce volume much greater than the Planck volume. The evolution of such states turns out to be in excellent agreement with the effective spacetime description derived from an effective Hamiltonian [21–23]. In this manuscript, we have extended the results on the quantum bounce to states which are very widely spread and which bounce much closer to the Planck volume. Investigations along these lines provide a very rigorous test for the validity of the effective dynamics. Numerical investigation of such states, on the other hand, present severe computational challenges which limit the types of states and the values of parameters for which a stable numerical evolution could be performed with previously used techniques. In order to overcome the computational limitations and study the numerical evolution of more general classes of states, we recently proposed a numerical scheme, Chimera, which proves to be a robust and efficient way to perform the simulations (see [29] for details). For example, as discussed in this paper, numerical simulations of very widely spread states, which would have taken extremely long time using conventional methods, can be performed in less than an hour on a workstation using the Chimera scheme.

In the present paper, we have used the Chimera scheme to study the LQC evolution of three different kinds of states and investigated the quantitative difference between the effective and LQC trajectories. We presented the results of a large number of simulations with various values of the relative volume dispersion $\Delta V/V$, spread in the field momentum $\Delta p_\phi$ and the field momentum $p_\phi$ at which the states are initially peaked. We found that the occurrence of the quantum bounce is a generic feature of all the different kinds of initial states we explored.
Figure 23. This figure shows the variation of $\delta$ with respect to the relative dispersion in volume $\Delta V/V$ for method-3 initial data. Panel (a) shows the full range of $\Delta V/V$, while panel (b) is a zoom into the small $\Delta V/V$ regime. It is clear from panel (a) that $\delta$ increases with increasing $\Delta V/V$ in the large $\Delta V/V$ regime. In panel (b) the dashed curve, connecting the turnaround points of the individual curves for different $p_\phi$ values, corresponds to $\Delta p_\phi = \tilde{\Delta} p_\phi = (3\pi p_\phi^2 G h^2)^{1/4}$.

This is in agreement with the results in [19], obtained via an exactly solvable model albeit for a slightly different quantum Hamiltonian constraint of the spatially flat isotropic model. We also found surprising quantitative results concerning the difference between the LQC and the corresponding effective trajectories for different types of initial data. As a generic feature, we found that the effective theory always overestimates the energy density at the bounce and underestimates the bounce volume, and that the effective theory remains a good approximation unless the states are widely spread. The energy density at the bounce always turns out to be bounded above by $\rho_{\max}$ predicted by the exactly solvable model [19] and the effective theory [equation (3.4)]. Our analysis also shows that the nature of departure of the effective theory
from LQC depends on the type of initial data, as well as the initial data parameters. To quantify these deviations we defined a quantity $\delta$ as the fractional difference in the bounce volume in the two descriptions. It turns out that $\delta$ can be large for small $p_0$, for which the bounce happens closer to the Planck volume. However, $\delta$ depends on the choice of initial state in a subtle way. For a fixed value of $\Delta V/V$, $\delta$ always increases with decreasing $p_0$ for method-1 and method-2 initial states, while for method-3 the behavior depends on whether $\Delta p_0$ is greater or smaller than a certain value. In the large $p_0$ regime, where $\delta$ is small, we found interesting non-monotonic variations of $\delta$ with $\Delta V/V$ and $\Delta p_0$. This behavior is similar to the non-monotonic dependence of $\Delta V/V$ on $\Delta p_0$ for the initial state. This was also noted in [31] for a different construction of initial state and a different quantum constraint. For all states and

\begin{figure}
\centering
\includegraphics[width=0.5\textwidth]{figure24}
\caption{This plot shows $\delta$ as function of $p_0$ for different values of $\Delta p_0$ for method-3 initial data. For a given $\Delta p_0$, $\delta$ is practically independent of $p^*_0$.}
\end{figure}

\begin{figure}
\centering
\includegraphics[width=0.5\textwidth]{figure25}
\caption{The behavior of the difference between the time of bounce in the LQC and the effective dynamical evolution for method-3 initial data. The curve is linear over the whole range, implying a power law relation between $(\phi_b - \phi_b^{(\text{eff})})$ and $\Delta p_0$. The slope of the curve is determined to be $\nu = -2.015 \pm 0.003$.}
\end{figure}
all initial conditions considered in this paper, our results are also in accordance with the study of the fluctuations of states across the bounce, and obey the corresponding triangle inequality \[30\] which places restrictions on the variation of the fluctuations across the bounce. Thus, in addition to strengthening some of the previous results on the quantum bounce, the results shown in this paper bring out some surprising features as well. For example, conventionally one would have expected that $\delta$ would continue to increase with increasing volume dispersion of the state. However, our analysis shows that this is not true in general.

It can be concluded from the numerical results obtained in this paper that the effective theory is in good agreement with the LQC evolution for states which are peaked at a classical macroscopic universe at late times. Notable differences between the effective and the LQC trajectories only occur for states with large relative fluctuations. In the derivation of the modified Friedmann equation in \[22\], such states lead to further corrections. Incorporating these corrections in the effective dynamics would lead to a better agreement with the numerical
simulations presented in this analysis. However, a more rigorous understanding is needed in the effective descriptions to gain insights on the way departures depend non-monotonically on fluctuations, and the way those departures are affected by the choice of the method of construction of the initial states. In addition, our results show that the role of fluctuations in the matter variable is as equally important as of the volume variable in understanding reliability of the effective dynamics\textsuperscript{14}. The matter and geometry fluctuations have non-trivial interdependence which can be different for different constructions of the initial states. Our results show that deciphering the reliability of effective descriptions in LQC is a subtle problem and an extra care is needed in generalizing conclusions from the effective descriptions.

Moreover, although the differences between the effective and the LQC trajectories for sharply peaked states corresponding to large field momenta are small, these differences depend on the parameters of the initial data in an unexpected fashion. Whether the effective theory becomes more or less reliable as one varies the parameters of the initial state, depends in a subtle way on the initial data parameters and the way the initial state is constructed. Note that in this paper, we have considered the effective description which is based on the embedding approach [21, 22]. The fate of the effective description based on the truncation approach [38] needs to be rigorously tested in a similar way, to make any reliable comparison between the two approaches to the effective theory and the loop quantum evolution.

Most of the phenomenological studies of various cosmological models and observational predictions of LQC are based on the effective description of LQC. The rigorous numerical simulations and the study of the differences between the effective and LQC trajectories, presented in this paper, open new avenues to understand the implications of the small departures between the effective theory and full quantum evolution on the observational signatures for widely spread states, e.g. along the lines of those computed in [26, 44]. For a better understanding of these modifications, one needs to extend the numerical studies to more general classes of models. This will be reported in upcoming works where we have used the Chimera scheme to study the numerical evolution of squeezed states and states with non-Gaussian waveforms for a massless scalar field [45] as well as more general matter models such as with an Ekpyrotic like potential [40]. These upcoming works along with the results presented in this paper provide a benchmark to numerically test the effective theory, and complement insights from the analytical derivation of the corrections to the effective equations for more general classes of states and matter models [23]. Availability of these corrections promises to provide a more rigorous understanding of the new physics resulting from the quantum bounce and its effects on cosmological perturbations.

Acknowledgments

We would like to thank Abhay Ashtekar, Steve Brandt, Frank Löffler and Miguel Megevand for useful discussions, and Martin Bojowald and Edward Wilson-Ewing for comments. This work is supported by a grant from John Templeton Foundation and by NSF grant PHYS1068743. The opinions expressed in this publication are those of authors and do not necessarily reflect the views of John Templeton Foundation. BG would also like to acknowledge support from the Coates Scholar Research Award and the Dissertation Year Fellowship of the Louisiana State University. This material is based upon work supported by HPC@LSU computing resources.

\textsuperscript{14} It will be interesting to investigate in detail the effects of matter fluctuations in the arguments presented in [42]. Study of these fluctuations is expected to yield more insights on the role of the infinite limit of the fiducial cell on the reliability of the effective dynamics.
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