Investigation of the stability of Hopfions in the two-component Ginzburg-Landau model

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We study the stability of Hopfions embedded in the Ginzburg-Landau (GL) model of two oppositely charged components. It has been shown by Babaev et al. [Phys. Rev. B 65, 100512 (2002)] that this model contains the Faddeev-Skyrme (FS) model, which is known to have topologically stable configurations with a given Hopf charge, the so-called Hopfions. Hopfions are typically formed from a unit-vector field that points to a fixed direction at spatial infinity and locally forms a knot with a soft core. The GL model, however, contains extra fields beyond the unit-vector field of the FS model and this can in principle change the fate of topologically non-trivial configurations. We investigate the stability of Hopfions in the two-component GL model both analytically (scaling) and numerically (first order dissipative dynamics). A number of initial states with different Hopf charges are studied; we also consider various different scalar potentials, including a singular one. In all the cases studied, we find that the Hopfions tend to shrink into a thin loop that is too close to a singular configuration for our numerical methods to investigate.

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I. INTRODUCTION

Topologically stable knots and other vortex-like structures have recently received wide interest within condensed matter physics. One reason for this is the continuous development of experimental techniques which now allow the production of vortices in various types of media. These include, e.g., Bose-Einstein condensates, superconductors, super-fluid $^3$He-A, and nematic liquid crystals. At the same time the increased capacity of supercomputers has made it possible to study these structures numerically. Thus, in the last decade there have been many numerical studies devoted to finding stable topologically non-trivial configurations in different physical systems, including, e.g., topological unknots, knots and vortices in the Faddeev-Skyrme (FS) model, vortexs in Bose-Einstein condensates, vortex atom lasers in a two-flavor Bose condensate, and vortices in superconductors, liquid helium, liquid metallic hydrogen, and possibly even in neutron stars.

There are various ways in which a vector field in nature can support vortices. For example a velocity field can form a vortex, e.g., in a hurricane. The position of the vortex is in the eye of the hurricane, where the velocity is zero. Such a vortex is not stable, because a velocity field can continuously change to zero everywhere.

In some special materials there can exist vector fields, associated to spin or other such property, that cannot vanish. Then it is possible to have vortices that are both non-singular and conserved, the conservation following from topological reasons. (One example that can be produced in a laboratory is the continuous unlocked vortex (CUV) in super-fluid $^3$He-A.) The prototype model that contains non-vanishing topological structures characterized by a Hopf charge was presented by L. Faddeev, and recently it has been shown numerically that this model does indeed contain stable topological solitons.

In this work we study numerically a system of two electro-magnetically coupled, oppositely charged Bose condensates described by the Ginzburg-Landau (GL) model. Babaev et al. studied this system in Ref. and argued that it should contain stable knotted vortex solitons with a non-zero Hopf charge. Here we present our results for the GL model, the main conclusion being that Hopfions in the GL model seem to be unstable.

The paper is organized as follows: In Section II we formulate the equations, discuss the possible potentials and present the initial states that are used in the computation. In Section III we present the discretization of the GL Lagrangian and the numerical method for finding stable minimum energy configurations. In Section IV we describe the results and finally in Section V we give some concluding remarks on the results obtained.

II. MODEL

A. Ginzburg-Landau model

The model describes two electro-magnetically coupled, oppositely charged Bose condensates, as given by the GL
Lagrangian density
\[
\mathcal{L} = \frac{\hbar^2}{2m_1} \left| (\nabla + i\frac{2e}{\hbar c} \vec{A}) \Psi_1 \right|^2 + \frac{\hbar^2}{2m_2} \left| (\nabla - i\frac{2e}{\hbar c} \vec{A}) \Psi_2 \right|^2 + V(\Psi_1, \Psi_2) + \frac{1}{2\mu_0} \vec{B}^2,
\]
where we have used SI units. In Eq. (1) \( \Psi_1 \) and \( \Psi_2 \) are the order parameters for the condensates, \( \vec{A} \) is the electromagnetic vector potential, \( \vec{B} \) the magnetic field, \( B = 4\gamma \nabla \times \vec{A}, \) and \( V \) is a potential.

Babaev et al.\cite{22} introduced new variables by setting
\[
\Psi_\alpha = \sqrt{2m_\alpha} \rho \chi_\alpha,
\]
where the new complex field \( \chi \) is normalized as
\[
|\chi_1|^2 + |\chi_2|^2 = 1,
\]
and therefore the real field \( \rho \) is given by
\[
\rho^2 = \frac{1}{2} \left( \frac{|\Psi_1|^2}{m_1} + \frac{|\Psi_2|^2}{m_2} \right).
\]
In terms of the new fields of (2) Eq. (1) becomes
\[
\mathcal{L} = \hbar^2 \rho^2 \left( |(\nabla + \frac{i2e}{\hbar c} \vec{A}) \chi_1|^2 + |(\nabla - \frac{i2e}{\hbar c} \vec{A}) \chi_2|^2 \right) + 2\hbar^2 (\nabla \rho)^2 + V(\chi_1, \chi_2, \rho^2) + \frac{1}{2\mu_0} \vec{B}^2.
\]

The Lagrangian (1) is invariant under the gauge transformation
\[
\begin{align*}
\Psi_1 &\to e^{-i\frac{2e}{\hbar c} \theta(x)} \Psi_1 \\
\Psi_2 &\to e^{i\frac{2e}{\hbar c} \theta(x)} \Psi_2 \\
A_\mu &\to A_\mu + c \partial_\mu \theta(x),
\end{align*}
\]
and the corresponding Noether current is
\[
J_k := \frac{i\hbar c}{m_1} (\Psi_1^* \partial_k \Psi_1 - \chi_1 \partial_k \chi_1^* - \chi_1^* \partial_k \chi_1 + \chi_2 \partial_k \chi_2^* + \chi_2^* \partial_k \chi_2) \\
- \frac{i\hbar c}{m_2} (\Psi_2^* \partial_k \Psi_2 - \chi_2 \partial_k \chi_2^* - \chi_2^* \partial_k \chi_2 + \chi_1 \partial_k \chi_1^* + \chi_1^* \partial_k \chi_1),
\]
which, using the Ampère’s law, must satisfy
\[
J = \nabla \times \vec{B} = \frac{1}{c} \nabla \times \nabla \times \vec{A}.
\]
When we use the new variables (2) here, we obtain
\[
J_k = 2e\hbar \rho^2 i(\chi_1^* \partial_k \chi_1 - \chi_1 \partial_k \chi_1^* - \chi_2^* \partial_k \chi_2 + \chi_2 \partial_k \chi_2^*) \\
- \frac{8e^2 \rho^2}{c} A_k
= 4e\hbar \rho^2 \left( \frac{j_k}{c} - \frac{2e}{c} A_k \right),
\]
which contains a new (non-gauge invariant) current
text
\[
jk = i(\chi_1^* \partial_k \chi_1 - \chi_1 \partial_k \chi_1^* - \chi_2^* \partial_k \chi_2 + \chi_2 \partial_k \chi_2).
\]
Later on we also use a gauge invariant vector field
\[
\vec{C} := \frac{1}{\hbar c \rho} \vec{J}.
\]
Next we define the unit vector field \( \vec{n} \) by
\[
\vec{n} = (\chi_1^* \chi_2) \sigma (\chi_1 \chi_2) \left( \frac{\chi_1 \chi_2 + \chi_1^* \chi_2^*}{|\chi_1|^2 - |\chi_2|^2} \right),
\]
where \( \sigma \) are the Pauli matrices. The inverse transformation is
\[
\begin{align*}
\chi_1 &= \frac{\alpha_1 - i\alpha_2}{\sqrt{2(1-\alpha_1\alpha_2)}} e^{i\alpha} \\
\chi_2 &= \frac{\sqrt{(1-\alpha_1)(1-\alpha_2)}}{\sqrt{2}} e^{-i\alpha},
\end{align*}
\]
where the phase must be chosen so that \( \chi_\alpha \) are continuous, see, e.g., (3). (This inverse transformation is not used in the numerical simulations.)

In order to show the similarities between the GL and FS models, we write the Lagrangian (5) in terms of \( \vec{n}, \rho \) and \( \vec{C}, \) after which the \( \vec{n} \) part of the result should be similar to the FS model, given by
\[
\mathcal{L}_{FS} = \frac{1}{2} \partial_k \rho \partial^k \rho^1 + g_{FS}(\vec{n} \cdot \partial_k \vec{n} \times \partial_l \vec{n}).
\]

After inverting Eqs. (9) and (11) to obtain \( \vec{A} \) in terms of \( \vec{C} \) and \( \vec{j} \) the kinetic part of Eq. (5) becomes
\[
\mathcal{L}_{\text{kinetic}} = \hbar^2 \rho^2 (|\nabla \chi_1|^2 + |\nabla \chi_2|^2 - \frac{1}{2} j^2) + 2\hbar^2 (\nabla \rho)^2 + \frac{\hbar^2}{16} c^2.
\]
Using Eqs. (12), (3) and (10) one finds that
\[
\partial_k \rho \partial^k \rho^1 = 4(|\nabla \chi_1|^2 + |\nabla \chi_2|^2) - j^2
\]
and therefore the first term in Eq. (15) corresponds to the first term in Eq. (14).

By direct substitution of Eqs. (9) and (11) into \( \vec{B} \), we also find that
\[
\vec{B} = \frac{1}{c} \nabla \times \vec{A} = \frac{\hbar}{2e} (\nabla \times \vec{j} - \frac{1}{2} \nabla \times \vec{C}),
\]
and again using Eqs. (12), (3) and (10), we get
\[
\frac{1}{2} \epsilon_{klm} \partial_k \vec{n} \times \partial_l \vec{n} = -\epsilon_{klm} \partial_k \vec{j},
\]
from which we can see that the \( \vec{B}^2 \) term in Eq. (15) contributes to the second term in Eq. (14).

Combining the above results we can write the Lagrangian (11) in the form
\[
\mathcal{L} = \frac{\hbar^2}{2m_1} (\partial_k \rho \partial^k \rho^1 + \hbar^2 (\nabla \rho)^2 + \frac{\hbar^2}{16} c^2 + V(\rho, n_k) + \frac{\hbar^2}{128\mu_0^2} \left[ \epsilon_{klm} (\vec{n} \cdot \partial_k \vec{n} \times \partial_l \vec{n} + \partial_k C_l) \right]^2,
\]
which is the form derived by Babaev et al.\cite{22}. The dynamical fields are now \( \rho, \vec{n} \) and \( \vec{C}. \) If \( \rho = \text{constant} \) and \( \vec{C} = 0, \) the GL model reduces to the FS model in Eq. (14). Since the FS model contains stable topological structures with non-trivial Hopf charge, one can hope that the GL model also contains similar structures. However, GL contains the additional fields \( \rho \) and \( \vec{C} \) in comparison to FS and the role of these new fields must be investigated.
B. Form of the potential

A typical and rather general quartic potential used in the GL model is

\[ V_0 (\Psi_1, \Psi_2) = \frac{1}{2}c_1|\Psi_1|^4 + \frac{1}{2}c_2|\Psi_2|^4 + c_3|\Psi_1|^2|\Psi_2|^2 + b_1|\Psi_1|^2 + b_2|\Psi_2|^2 + a_0. \]  

(20)

When \( \Psi_\alpha \) are expressed in terms of \( \rho \) and \( \vec{n} \) and the whole system is rescaled so that \( m_\alpha \to 1 \), we find

\[ |\Psi_1|^2 = \rho^2(1 + n_3), \quad |\Psi_2|^2 = \rho^2(1 - n_3), \]  

(21)

and then the potential \((20)\) reads

\[ V_0 (\rho^2, n_3) = \rho^4 [n_3^2 \left( \frac{1}{2}(c_1 + c_2) - c_3 \right) + n_3(c_1 - c_2) + \frac{1}{2}(c_1 + c_2 + c_3)] + \rho^2 [n_3(b_1 - b_2) + b_1 + b_2] + a_0. \]  

(22)

One important aspect in choosing the potential is that at infinity the fields will settle to the minimum of the potential. Furthermore, in order to define the Hopf charge it is necessary that the \( \vec{n} \) field points to the same direction far away, otherwise we cannot compactify the 3D-space.

For particular parameter values the extrema are obtained for particular values of \( n_3 \) and \( \rho^2 \):

\[ \begin{align*}
 n_3 &= \frac{b_1(c_2 + c_3) - b_2(c_1 + c_3)}{b_1(c_2 - c_3) + b_2(c_1 - c_3)}, \\
 \rho^2 &= -\frac{b_1(c_2 - c_3) + b_2(c_1 - c_3)}{2(c_1c_2 - c_3^2)}. 
\end{align*} \]  

(23)

This is a minimum, if \( c_1c_2 > c_3^2 \). Note that the above values do not necessarily fall within the allowed values for \( \rho^2 \) and \( n_3 \), i.e., \( \rho^2 > 0 \) and \( |n_3| \leq 1 \), in which case the extrema are on the boundaries of the allowed values.

From physical arguments the following special case is relevant:

\[ V_1 (\Psi_1, \Psi_2) = \lambda \left( (|\Psi_1|^2 - 1)^2 + (|\Psi_2|^2 - 1)^2 \right). \]  

(25)

This breaks \( O(3) \) to \( O(2) \) and corresponds to two independently conserved condensates. It has a minimum at \( n_3 = 0, \rho^2 = 1 \).

Another physically relevant potential is

\[ V_2 (\Psi_1, \Psi_2) = \lambda \left( (|\Psi_1|^2 - 1)^2 + (|\Psi_2|^2 - 1)^2 \right) + c|\Psi_1\Psi_2^* - \Psi_2\Psi_1^*| + a_0, \]  

(26)

which breaks \( O(3) \) completely. This corresponds to multi-band superconductors with inter-band Josephson effect. In terms of \( \vec{n} \) this potential is given (using Eq. [13])

\[ V_2 = \lambda \frac{n_3^2}{2} + \text{Im}(n_2 - i\text{n}1)e^{i\alpha} + b_0. \]  

(27)

The minimum of this potential is located at \( \rho^2 = 1/2, n_3 = \sqrt{1 - c^2}/\lambda \), while the specific values of \( n_1 \) and \( n_2 \) also depend on \( \alpha \), which is related to the phase of \( \Psi \).

From the point of view of Hopf-charge conservation, the possibility of \( \rho = 0 \) is problematic, even if it happens locally, since then the field \( \vec{n} \) is not defined. However, it has been argued\(^{20}\) that quantum effects ensure that \( \rho \neq 0 \) everywhere. In classical field theories, like the present one, such expected quantum effects can be included through effective potentials. Since the main purpose of this effective potential is to guarantee that \( \rho \neq 0 \), its exact form is not so important. One such potential, which we will use later, is

\[ V_{\text{eff}} (\Psi_1, \Psi_2) = \frac{1}{2} \lambda \left( |\Psi_1|^2 + |\Psi_2|^2 - r_0^2 \right)^2 + 2\gamma \left( |\Psi_1|^2 + |\Psi_2|^2 \right)^{-1} + a_0. \]  

(28)

The constants \( r_0 \) and \( a_0 \) are determined by requiring that the minimum of the potential is at \( \rho^2 = 1 \) with a value of 0, yielding \( r_0^2 = 2 - \gamma/\lambda \) and \( a_0 = -\gamma - \frac{2}{\lambda} \).

C. Initial states of Hopf invariant \( \mathcal{Q} \)

We are interested in the minimum energy configurations of topologically distinct configurations of the field \( \chi \), or by Hopf-map, \( \vec{n} \), related to the complex physical fields \( \Psi_\alpha \) through Eqs. [13] and [14]. From the point of view of \( \vec{n} \) it is only necessary that \( \lim_{|\vec{n}|\to\infty} \vec{n} = \bar{n}_\infty \) is the same in all directions. From this it follows that we can compactify \( \mathbb{R}^3 \to S^3 \) and then \( \vec{n} \) becomes a map \( S^3 \to S^2 \) with homotopy group \( \pi_3 (S^2) = \mathbb{Z} \), characterized by the Hopf charge.

Therefore, we have to create a configuration with \( \Psi : \mathbb{R}^3 \to \mathbb{C}^2 \) and \( \tilde{A} : \mathbb{R}^3 \to \mathbb{R}^3 \), such that \( \vec{n} \) has the desired property mentioned above. For \( \Psi_\alpha \) this implies \( |\Psi_1|^2 + |\Psi_2|^2 \neq 0 \) everywhere and \( \lim_{|\vec{x}|\to\infty} \Psi_\alpha \) are (independent) constants. The first condition is also necessary for defining the field \( \chi \) and if the second condition is also satisfied \( \chi \) becomes a map \( S^3 \to S^3 \).

The method of constructing a configuration with a desired Hopf charge has been investigated by Aranyu et al.\(^{31}\). They used the fact that for any function \( \phi : S^3 \to S^3 \) combined with the Hopf map \( h : S^3 \to S^2 \)

\[ h (\phi_1, \phi_2, \phi_3, \phi_4) = \left( \frac{2(\phi_1\phi_3 - \phi_2\phi_4)}{\phi_1^2 + \phi_2^2 - \phi_3^2 - \phi_4^2}, \frac{2(\phi_2\phi_3 + \phi_1\phi_4)}{\phi_1^2 + \phi_2^2 - \phi_3^2 - \phi_4^2}, \right) \]  

(29)

the Hopf charge of \( h \circ \phi : S^3 \to S^2 \) equals the degree of the map \( \phi : S^3 \to S^3 \) (the degree is the \( n \)-dimensional generalization of the 1-dimensional degree, also known as the winding number).
For an explicit construction one uses the toroidal coordinates \((\eta, \xi, \varphi)\) of \(\mathbb{R}^3\) defined by
\[
\begin{align*}
x_1 &= \frac{\sinh(\eta) \cos(\varphi)}{\Delta}, \quad x_2 = \frac{\sinh(\eta) \sin(\varphi)}{\Delta}, \\
x_3 &= \frac{\sin(\xi)}{\Delta}, \quad \Delta = \cosh(\eta) - \cos(\xi).
\end{align*}
\] (30)

In these coordinates the core is at \(\eta = \infty\), while the z-axis and spatial infinity are at \(\eta = 0\).

Next take any monotonic function \(g: [0, \infty) \rightarrow [0, 1]\), (or \([-1, 0]\)) choose \(p, q \in \mathbb{Z}\) and define the map \(\phi: \mathbb{R}^3 \rightarrow \mathbb{S}^1\) by
\[
\phi = \left( g(\eta) \cos(p\xi), g(\eta) \sin(p\xi), \sqrt{1 - g(\eta)^2 \cos(q\varphi)}, \sqrt{1 - g(\eta)^2 \sin(q\varphi)} \right).
\] (31)

If furthermore \(g\) is such that \(g^2(\infty) - g^2(0) = \pm 1\), then the combined map, \(h \circ \phi\), has the Hopf invariant
\[
H(h \circ \phi) = \pm pq.
\] (32)

For details, see Ref.\(^\text{21}\). Using (31) we now identify
\[
\begin{align*}
\chi_1 &:= \phi_1 + i \phi_2 = g(\eta) e^{ip\xi}, \\
\chi_2 &:= \phi_3 + i \phi_4 = \sqrt{1 - g(\eta)^2} e^{iq\varphi}.
\end{align*}
\] (33a)
(33b)

At the z-axis \(\chi_2\) looks like a \(\varphi\)-vortex, and therefore for continuity we add the further requirement that \(g(0) = \pm 1\). Similarly, around the core (located at \(\eta = \infty\)) we have a \(\xi\)-vortex, and therefore we demand that \(g(\infty) = 0\).

From Eqs. (12) and (33) we obtain
\[
\vec{n}(\vec{x}) = \left( \frac{2g(\eta) \sqrt{1 - g^2(\eta)^2 \cos(p\xi + q\varphi)}}{2g^2(\eta) - 1}, \frac{-2g(\eta) \sqrt{1 - g^2(\eta)^2 \sin(p\xi + q\varphi)}}{2g^2(\eta) - 1} \right).
\] (34)

Finally, to close the loop, the \(\chi\) of (33) can be recovered from (13) using (34) and choosing \(\alpha = -q\varphi\).

Note that since \(\eta = 0\) at infinity and \(g(0)^2 = 1\) the above construction implies \(\vec{n}_\infty := \lim_{|\vec{x}| \to \infty} \vec{n} = (0, 0, 1)\).

Inverting the toroidal coordinates defined in Eq. (30) enables us to express Eq. (33) in the Cartesian coordinates. In addition, denoting \(r^2 = x_1^2 + x_2^2 + x_3^2\) and choosing \(\rho = 1\), \(g(\eta) = 1/\cosh(\eta)\) we obtain
\[
\begin{align*}
\Psi_1(\vec{x}) &= \sqrt{\frac{r^2 - 1}{r^2 + 1}} \left( \frac{1}{\sqrt{r^2 - 2r + 2}} \right)^p, \\
\Psi_2(\vec{x}) &= \frac{2}{r^2 + 1} \left( \frac{x_1 + 1}{\sqrt{r^2 + 1}} \right)^q.
\end{align*}
\] (35a)
(35b)

This is the formula used for the initial configurations of \(\Psi_\alpha\) for our numerical computations. The state indeed has the correct Hopf invariant, which, in the case of \(p = q = 1\) can be seen by finding the preimages of \(\Psi_\alpha = 0\), which correspond to preimages of \(n_3 = \pm 1\). These form two closed loops, namely \(r = 1, x_3 = 0\) for \(\Psi_1 = 0\) and the z-axis, which in the compactified space \(\mathbb{S}^3\) is actually closed.

Fixing \(\vec{n}\) or \(\chi_i\) does not say anything about the magnitude of \(\Psi\). From energy consideration we must choose \(\lim_{|\vec{x}| \to \infty} \Psi_\alpha\) to be one of the minima of the potential and this fixes a preferred value for \(\rho\). We use this preferred value for all \(\vec{x}\) when constructing the initial configurations.

### III. NUMERICs

In this section we will describe the discretization of the model and the method of the minimization of the Lagrangian (11). We will also compare the GL to FS model and present some test calculations and the parameters as well as coupling constants for the simulations.

It is important to note that all our simulations are done with the fields \(\Psi_\alpha\) only, the fields \(\vec{n}, \rho\) and \(\vec{C}\) are never used in any computation, they are only used in the analysis of the results.

It is perhaps useful to mention once again that the change of variables in Eqs. (24) is not reversible whenever \(\rho(\vec{x}) = 0\). There is, however, no guarantee that \(\rho\) stays non-zero in numerical simulations of the physical fields \(\Psi_\alpha\) and \(\vec{A}\), even if the minimum of the potential is at a non-zero value of \(\rho\). The situation when \(\rho = 0\) locally can imply breakdown in topology and therefore in our numerical simulations we have monitored the changes in the global minimum value of \(\rho\).

Topology can also break if the topological structure shrinks smaller than the lattice unit length. In order to be aware of this possibility we have monitored the global minimum of the dot product of nearest-neighbor \(\vec{n}\)-vectors: when the global minimum becomes negative the solution is probably too coarse, or the configuration has a genuine singularity. When this has happened in the simulations, we have repeated the simulation with ever increasing lattice size until the computing resources were exhausted. This strongly points to a singular configuration.

For simplicity we have used the rescaled \(|\Psi_\alpha|^2/m_\alpha \rightarrow \{\Psi_\alpha|^2\}_\alpha\) Lagrangian and natural units \((c = \hbar = 1)\) throughout our numerical work. With these choices, the Lagrangian density used in all our numerical simulations becomes
\[
\mathcal{L} = \frac{1}{2} \left( \nabla + ig \vec{A} \right) \Psi_1^2 + \frac{1}{2} \left( \nabla - ig \vec{A} \right) \Psi_2^2 + \frac{1}{2} g^2 |\vec{A}|^2 + V(\Psi_{1,2}).
\] (36)

This can be further scaled by \(\vec{A} \rightarrow \frac{1}{\sigma} \vec{A}\), which reveals the fact that the only relevant parameter is \(g^2/g_f = 4\mu_0e^2\).

The values used in numerical simulations are \(g_1 = 1\) and \(g_f \in \{0.01, 1, 2, 100\}\); these amount to a particular choices of units for \(\mu_0\) and \(e\).

In studies of the FS model, we have always assumed that \(\lim_{|\vec{x}| \to \infty} n_3 = 1\), but some of the present potentials, e.g., \(\hat{V}_1\), do not have that as a minimum. In this case we may assume that \(\lim_{|\vec{x}| \to \infty} n_2 = 1\), but this
can be transformed to \( n_3 = 1 \) by a global rotation in the configuration space, which takes \( n_3 \rightarrow n_2 \) and \( n_2 \rightarrow -n_3 \). The same effect can be achieved by using new fields defined by

\[
\Psi'_1 = \frac{1}{\sqrt{2}}(\Psi_1 + i\Psi_2), \quad \Psi'_2 = \frac{1}{\sqrt{2}}(\Psi_1 - i\Psi_2),
\]

changing \( V_1 \) of (25) to

\[
V_{1\text{rot}} = \lambda\left((|\Psi'_1|^2 + |\Psi'_2|^2 - 2)\right)^2 \\
- \left((\Psi_1\Psi_2 - \Psi'_1\Psi'_2)^2\right). \tag{38}
\]

The different potential terms also contain various parameters. For \( V_0 \), we have always used \( 4c_1 = 4c_2 = 2c_3 = -b_1 = -b_2 = a_0 \), which, denoting \( \lambda \equiv \frac{1}{2}c_1 \), enables us to write \( V_0 = \lambda(|\Psi|^2 + |\Psi|^2 - 2)^2 \). In the numerical simulations, we have used potentials \( V_0, V_{1\text{rot}} \) and \( V_{\text{eff}} \) with \( \lambda \in \{0, 1, 4, 100, 1000\} \).

### A. Discretization

The system has been discretized on a cubic rectangular lattice (indexed as \((s, u, v)\)) with periodic boundary conditions. Our model can be considered as a two-component version of the time-independent, ordinary Abelian U(1) Higgs model, which has long since been discretized for lattice simulations in quantum field theory (for example, see Refs. 22 and 23 and the references therein). The main point in that context is to discretize the fields so that gauge invariance is preserved. From Eq. (9) we see that if we use the forward discretization of the derivatives, the gauge transformation of \( A_k \) has to be discretized as follows:

\[
A_{1|s,u,v} \rightarrow A_{1|s,u,v} + \frac{\xi}{a}(\theta_{s+1,u,v} - \theta_{s,u,v}), \tag{39a}
\]

\[
A_{2|s,u,v} \rightarrow A_{2|s,u,v} + \frac{\xi}{a}(\theta_{s,u+1,v} - \theta_{s,u,v}), \tag{39b}
\]

\[
A_{3|s,u,v} \rightarrow A_{3|s,u,v} + \frac{\xi}{a}(\theta_{s,u,v+1} - \theta_{s,u,v}), \tag{39c}
\]

where \( a \) is the lattice parameter. Thus, \( A_k \) should be considered as living on the link between two lattice points parallel to the coordinate axis \( k \). Combining this discretization with the way the \( \Psi_\alpha \) transforms in Eq. (9) it follows that the following combination

\[
\Psi_{1|s+1,u,v} \Psi_{1|s,u,v} e^{-i\kappa A_{1|s,u,v}} \\
+ \Psi_{1|s+1,u,v} \Psi_{1|s,u,v} e^{i\kappa A_{1|s,u,v}} \\
- \Psi_{1|s+1,u,v} \Psi_{1|s,u,v} - \Psi_{1|s,u,v} \Psi_{1|s+1,u,v}, \tag{40}
\]

where \( \kappa = \frac{2\pi}{a} \), is gauge invariant. If we calculate its continuum limit as \( a \rightarrow 0 \) by expanding in \( a \) (e.g., \( \Psi_{1|s+1,u,v} = \Psi_{1}(x + a, y, z) \)) we obtain \(-a^2(\partial_1 + i\kappa A_1)\Psi_{1}^2 + O(a^2)\). For \( \Psi_2 \) with \( A_1 \) we use a similar expression with \( \kappa \rightarrow -\kappa \). Finally, for the other components of \( \vec{A} \) we use corresponding shifts, as illustrated in Eqs. (39) and (10).

For the discretization of \( \vec{B}^2 \) we use the expression

\[
e^{iF_{12|s,u,v}} + e^{-iF_{12|s,u,v}} + e^{iF_{23|s,u,v}} + e^{-iF_{23|s,u,v}} \\
+ e^{iF_{31|s,u,v}} + e^{-iF_{31|s,u,v}} - 6, \tag{41}
\]

where, for example,

\[
F_{12|s,u,v} = A_{1,s,u+1,v} - A_{1,s,u,v} - A_{2,s+1,u,v} + A_{2,s,u,v}. \tag{42}
\]

The \( F_{kl} \) are gauge invariant under Eq. (39), and the continuum limit of Eq. (11) with Eq. (12) is \(-a^2(\nabla \times \vec{A})^2 + O(a^3)\).

For the potential we use \( a^2V(\Psi_{\alpha|s,u,v}) \). The discretized Lagrangian is the sum of all the above terms multiplied by \(-a/2\), and then the resulting sum has the continuum limit of Eq. (36). In practice we use the cubic lattices of sizes of \( 60^3 \ldots 960^3 \). Of these just the sizes \( 240^3, 480^3 \) and \( 720^3 \) are used in actual simulations, the remaining sizes used only to verify the code, results and discretization.

The minimization of the Lagrangian has been done using the steepest descent method and the Fletcher-Reeves variant of the conjugate gradient method. The gradients needed have been calculated symbolically from the discretized Lagrangian.

### B. Initial states

In order to start the simulations we have to generate initial states with specified Hopf invariants. These initial configurations have been made using Eq. (22) and choosing \( \rho \) such that \( V(\Psi_\infty) = 0 \). Three different values of Hopf invariant were used: 1, 2 and 4. However, this leaves \( \vec{A} \) undetermined. The topology of the system is unaffected by \( \vec{A} \), so in principle, we can choose any configuration for it. We have used several different initial configurations for \( \vec{A} \); for all the results presented here, we have used two choices. The first one is defined by the condition \( \vec{C} = 0 \Rightarrow \vec{A} = \frac{\vec{B}}{2}, \) by Eqs. (10) and (11). This choice enables us to test the validity of the program by comparing the energies of the initial states with those obtained for the pure FS model. The second initial configuration has been constructed by solving Ampère’s law for \( \vec{A} \) with fixed \( \Psi_\alpha \). Note that since we use gradient methods to minimize the energy, we already have encoded Ampère’s law in our program: the gradients of the Lagrangian with respect to \( \vec{A}_k \) indeed yield Ampère’s law.

As a further test of discretization we used the same initial configurations in lattices of different sizes and with different lattice constants. With lattice sizes of \( 240^3 \) and above the energies were within 1% of each other.
C. Comparison with previous FS studies

We have also compared the new calculations with those presented in Ref. 13. By using an initial configuration, where \( \rho \equiv 1 \), \( \dot{C} \equiv 0 \) and \( V \equiv 0 \), the GL model (19) reduces to the FS model (14). In particular, the magnetic term becomes \( E_{M_B} = \frac{\hbar^2}{2e} \epsilon_{klm} (\vec{n} \cdot \partial_l \vec{n} \times \partial_m \vec{n})^2 \), while in the FS model we have \( E_{TFS} = \frac{1}{2} g_{FS} (\vec{n} \cdot \partial_i \vec{n} \times \partial_j \vec{n})^2 \). If we set \( c = h = 1 \), \( e = \frac{1}{2} \), \( \mu_0 = 1 \) and \( g_{FS} = \frac{1}{2} \) (the value most commonly used in our earlier work), we have \( E_{TFS} = 2E_{M_B} \). We created the same initial configuration with our old and new codes and found that the energies of the initial states agree to within 1% with a lattice of 120³.

During the calculations we have monitored the topology of the system. As we saw in Sec. 13 the conservation of topology is not guaranteed whenever \( \rho \rightarrow 0 \). Therefore we have followed the value of the global minimum of \( \rho^2 \). In keeping with our earlier work, we have also monitored the dot products of \( \vec{n}(\vec{x}) \cdot \vec{n}(\vec{x} + \vec{\mu}) \), (where \( \vec{\mu} \in \{(1,0,0),(0,1,0),(0,0,1)\} \)). When the global minimum of this dot product approaches zero or becomes even negative it indicates possible breakdown of continuity and therefore of topology during the simulation.

We have also calculated the value of the Hopf invariant directly from the field configuration. Using the basic differential geometric result of \( H(\dot{h} \circ \chi \equiv \vec{n}) = \deg \chi \), we can even use our computational field variables. Since for any map \( f \in C^\infty(S^3 \rightarrow S^3) \),

\[
\deg f = \frac{1}{(2\pi)^3} \int_{S^3} \epsilon_{\mu \nu \rho} f^\mu df^\nu \wedge df^\rho \wedge df^\rho,
\]

we can compute the value of the Hopf invariant by numerical integration. The Hopf invariant can also be determined by visual inspection of the linking numbers of the preimages.

IV. RESULTS AND DISCUSSION

A necessary condition for any stable static localized structure is its stability against scaling. Let us assume, for now, that \( \rho \neq 0 \) everywhere. Then the simple scaling argument of Derrick 34 gives a necessary condition for the stability against scaling. Consider the behavior of the energy under scaling \( x \rightarrow \lambda x \). For the FS model in 3D the volume element of the energy integral behaves like \( \lambda^{-3} \) while in the integrand (14) the first term scales as \( \lambda^{-2} \) and the second one as \( \lambda^{-4} \). In the integral the two terms then have opposite scaling behavior and if each is positive definite, one cannot scale the energy to zero. Furthermore, by the virial theorem the preferred scaling is the one where the two terms have equal value. In fact one can show 35 that in the FS model the energy is bounded from below by the Hopf charge: \( E > c|Q|^{3/4} \). This has also been confirmed numerically. 10,11,12,33

In the present model the scaling stability hinges on the behavior of the first and last terms in Eq. (19). These terms are positive definite and their scaling properties are opposite and thus, as long as they are both non-zero, the system is stable. The last term, however, equals the term \( \vec{B}^2 \) in Eq. (1) and could, in principle, vanish. Indeed, if the \( \dot{C} \) field obtained such a value that

\[
\epsilon_{klm} \partial_i C_m = -\epsilon_{klm} \vec{n} \cdot \partial_i \vec{n} \times \partial_m \vec{n},
\]

then the last term would vanish, rendering the system unstable against scaling. One could say that topological charge leaks from \( \vec{n} \) to \( \dot{C} \), so that \( \vec{B} \rightarrow 0 \).

We have observed the above scaling instability in direct minimizations of all fields. The minimization does not exactly follow the route of a uniform scaling, but we have observed that the torus-like un-knot shrinks into a thin tube, and the value of the integrand in Eq. (19), which contributes to the topological charge, concentrates tightly in a region near the vortex core. Eventually the nearest neighbor values in the corresponding \( \vec{n} \)'s are anti-parallel, which in effect means that the vortex configuration has shrunk to a size less than the lattice spacing. This is accompanied with \( \rho \) approaching zero, and we have verified that \( \rho \) approaches zero at the same location where the neighboring \( \vec{n} \)’s are anti-parallel. At a place where \( \rho = 0 \) the field \( \vec{n} \) can no longer be determined from \( \Psi \), and the topological structure breaks, after which the system rapidly goes to a vacuum state. We have observed this scenario for various different potential forms and strengths.

Two steps of the process of topology breakdown are illustrated in Fig. 14 where we have plotted two distinct preimages of \( \vec{n} \), \( n_2 = 1 \) and \( n_2 = -1 \) and the isosurface \( n_3 = 0 \) on which the tubes lie: the colours (gray-scale) of the isosurface describe longitudes. Initially (left image) the preimages are spaced 90 degrees apart on the isosurface \( n_3 = 0 \), but during the minimization, on the region of \( xy \)-plane inside the torus core, the preimages deform so that eventually there are no lattice points between the preimages of \( n_2 = 1 \) and \( n_2 = -1 \). This happens even with \( V_{eff} \) (in which case \( \rho > 0 \) everywhere) and is not accompanied by a shrinking torus; the torus deforms somewhat but the radius of the inner intersection of \( n_3 = 0 \) and \( xy \)-plane stays approximately constant. The two preimages eventually touch each other and when they drift apart again (right image), they have become reconnected so that the topology is trivial. After this, there is nothing to prevent, the energy from dropping to zero.

It should be noted, however, that for any non-trivial configuration \( \Psi_\alpha \), Eq. (14) violates Ampere’s law, Eq. (8), because by using Eqs. (15), (17), and (8), Eq. (44) implies \( \vec{B} = 0 \) and \( \nabla \times \vec{B} = \vec{J} = 0 \), which contradicts the topological non-triviality of \( \Psi_\alpha \). Thus any non-trivial configuration, where Eq. (44) holds, is unphysical. It can then be argued, that although the scaling route leads to instability, it does so via non-physical states and
thus does not imply the instability of physically relevant states.

In order to stay within physically relevant states during minimization, we have used the following procedure: After every minimization step applied to the $\Psi$ fields we check whether Ampère’s law is satisfied. At the beginning of a simulation the $\Psi$ fields are generated from analytical formulae, such as $\Psi = (1/2\alpha \vec{n}_1 \vec{A})$, which provide the proper topological charge. Then Ampère’s law is solved for this initial state (using conjugate gradients) until it is “sufficiently accurate”. After this initial step we take conjugate gradient iteration steps for all fields until the convergence criterion for $\vec{A}$ is no longer satisfied, at which point we continue only with $\vec{A}$ and Ampère’s law until it is satisfied to the desired accuracy. This is repeated until the whole iteration has converged. There are various methods to determine when the solution is “sufficiently accurate”. Since we use a gradient-based method to solve Ampère’s law, it is natural to determine the accuracy of the solution using the gradients. To accomplish this, we observe the absolute values of the gradients affecting $\vec{A}$ and note the maximum of these values at each iteration and the initial state before any iterations are made. We then compare these maxima with the maximum found at the initial state; the solution is considered “sufficiently accurate” if the ratio of the current maximum to the initial maximum is below 0.0001.

The result of the minimization process is always the same: a singular configuration of $\Psi_1$, where the region contributing to the topological charge has shrunk to a singular line. At the same time $\vec{B}$ field is generated in the hole of the torus. This can be seen from Fig. 2 which describes states before and just after topology breakdown. In particular one can see how the $\vec{B}$ is formed mostly on a smaller ring in the torus hole and how the initial maximum of $\Psi_1$ at origin vanishes while its toroidal minimum becomes disk shaped. The topology is broken in exactly the same way as in the unconstrained case.

The exact type of singularity and the process which leads to its formation, depends on the potential. For $V_0$ and $V_{\text{eff}}$, the small radius of any initial torus-type isosurface of $n_3$ shrinks without limit, snaring the region of topological interest into a singular loop. For $V_{\text{nat}}$, the process is the same unless the potential is very strong, in which case the region where $n_3 \neq 0$, shrinks to a surface.

This process is ultimately a scaling instability, albeit a different one from the one considered at the beginning of this Section. Now, the scaling is not global, but only shrink the smaller circle of the torus. The terms involving derivatives of $\Psi_1$ grow without limit during the shrinking process, but only on a loop or a surface, allowing still the total energy to decrease.

In every case the shrinking continues until the discreteness of the computational lattice eventually breaks the topology. We have tried to follow the development of this singularity by increasing lattice density. In each case a singularity is reached, but in principle the shrinking can stop at some still smaller scale determined by the dimensional parameters of the system (such as penetration length, strength of the potential, etc.).

V. CONCLUSIONS

We have investigated numerically the two-component Ginzburg-Landau model using as initial states torus unknots, with non-vanishing Hopf change $\mathcal{Q}$; this brings a topological structure into the system. We have used different types of potentials depending on both order parameters $\Psi_1$ of the system. In all cases the initial torus tube shrinks into a thin loop and becomes untrackable in our computational discrete lattice: the discreteness allows the topological structure of the system to disappear eventually, contrary to the case of the Faddeev-Skyrme model. The topological stability in the FS model is due to the fact that the kinetic and topological terms are non-vanishing and have an opposite behavior in the scaling.

In the GL model, the term corresponding to the FS topological term is the magnetic field term which contributes to the unstability of the system. In a direct minimization of the fields, the topological charge leaks from $\vec{n}$ into $\vec{C}$ allowing eventually Derrick-type instability, while in minimizations respecting Ampère’s law the process is milder, but nevertheless leads to singularity.

Thus, it seems that the two-component GL model does not support stable topological structures having a non-trivial conserved Hopf invariant, due to this scaling instability. It is still possible that in the GL-system the natural size of a stable un-knot is much smaller than in the FS-model, or that it is only stable for suitably strong potentials. It is also possible to stabilize the topological structures by adding a suitable term in the Lagrangian. One such term has been introduced in Ref. There are questions that we will study further.
FIG. 2: Snapshots of a $Q = 2$ system before ((a) and (c)) and just after ((b) and (d)) the topology breakdown. Cross-sections of the system have been taken in $xy$-plane ((a) and (b)) and $xz$-plane ((c) and (d)). In all four panels, the upper two plots describe the densities $|\Psi_1|^2$ (left) and $|\Psi_2|^2$ (right) on which the phases of the condensates are superimposed as contours spaced $\pi/3$ apart; the lower two plots of each panel describe the energy densities $\frac{1}{2}g_1|\vec{B}|^2$ (left) and $E_{\text{total}} - \frac{1}{2}g_1|\vec{B}|^2$ (right). The jagged line in the $xy$-cross-sections corresponds to $\phi = \pi$; the jaggedness itself is an imaging artifact caused by the discreteness of the data used to produce the image.
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