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Topical Review

Theoretical estimates of maximum fields in superconducting resonant radio frequency cavities: stability theory, disorder, and laminates

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Abstract
Theoretical limits to the performance of superconductors in high magnetic fields parallel to their surfaces are of key relevance to current and future accelerating cavities, especially those made of new higher-Tc materials such as Nb₃Sn, NbN, and MgB₂. Indeed, beyond the so-called superheating field $H_{sh}$, flux will spontaneously penetrate even a perfect superconducting surface and ruin the performance. We present intuitive arguments and simple estimates for $H_{sh}$, and combine them with our previous rigorous calculations, which we summarize. We briefly discuss experimental measurements of the superheating field, comparing to our estimates. We explore the effects of materials anisotropy and the danger of disorder in nucleating vortex entry. Will we need to control surface orientation in the layered compound MgB₂? Can we estimate theoretically whether dirt and defects make these new materials fundamentally more challenging to optimize than niobium? Finally, we discuss and analyze recent proposals to use thin superconducting layers or laminates to enhance the performance of superconducting cavities. Flux entering a laminate can lead to so-called pancake vortices; we consider the physics of the dislocation motion and potential re-annihilation or stabilization of these vortices after their entry.

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Keywords: superheating field, superconducting radio frequency cavities, flux penetration, disordered nucleation

(Some figures may appear in colour only in the online journal)

1. Introduction

To transfer energy to beams of charged particles, accelerators frequently use superconducting radio-frequency (SRF)
cavities, devices that are capable of sustaining large amplitude electromagnetic fields with relatively small input power. The energy gain of a beam traversing a cavity is determined by the electric field amplitude along its path—a larger amplitude can reduce the number of cavities required to reach a given energy. This is especially important in high energy accelerators, which call for as many as tens of thousands of cavities [1]. It is therefore of interest to understand the mechanisms that fundamentally limit the accelerating electric field. For state-of-the-art SRF cavities that have been carefully prepared to prevent non-fundamental degradation processes such as field emission [2, 3] and multipacting [4], studies show that the limit is not the electric field, but rather the interaction of the magnetic field with the superconducting material of the cavity walls. The fundamental limit to acceleration in SRF cavities is the superheating field $H_{sb}$, introduced in section 2.

This article will cover ideas, methods, and results revolving around the superheating field and its dependence on the superconductor—materials properties, anisotropy, defects and disorder, and laminates. The ideas and methods are primarily gleaned from the broader condensed matter community. In section 2 we review computations of $H_{sb}$ for clean systems using field theories from the 1950s derived for pure superconductors near their transition temperature [5]; in section 4.1 we draw from more sophisticated theories from the 1960s to calculate $H_{sb}$ at all temperatures [6], and discuss the future need to use these historical theories to incorporate effects of strong coupling and electronic structure [7] in new materials. In section 4.2 we review the use of these methods to address the electronic anisotropy of some of the new materials. In section 4.3 we introduce an illustrative calculation of the effects of disorder using tools and methods developed in the 60s for disordered systems [8, 9] and nucleation theory [10, 11], providing reassurance that new materials will likely not be far more sensitive to flaws and dirt. Finally, in section 5 we investigate the properties of superconducting laminates, by drawing from work from the 90s on the dynamics of ‘pancake vortices’ in certain layered high-temperature superconductors [12] (particularly BSCCO, Bi$_2$Sr$_2$Ca$_n$CuO$_{2n+4+x}$).

We frankly have two goals for this article. As discussed above, we wish to provide an introduction for the accelerator community into tools and methods from the broader condensed matter community that can help interpret current experimental challenges and guide plans for future research in optimizing materials properties for SRF cavities. But conversely, we want to provide a window for the broader condensed matter theory community into the remarkable frontiers of field, frequency, and materials preparation being explored by the SRF community. We invite their participation in melding 21st century materials-by-design tools from electronic structure theory with 20th century field theories of superconductivity, bridging the scales to address current technological challenges in the accelerator field. (Full disclosure: this article was supported in part by the Center for Bright Beams, an NSF Science and Technology Center whose mission is precisely to bring the accelerator community together with outside experts in physical chemistry, materials science, condensed matter physics, plasma physics and mathematics.)

1.1. Basic facts about superconductors: type I and II, $H_c$, $H_{c1}$, and $H_{c2}$

Normal conducting metals, such as copper, are not viable as radio-frequency cavities for long-pulse high-gradient applications. Due to their high surface resistance, these cavities dissipate too much power on the walls, which can result in melting, among other structural problems, if they are not sufficiently cooled. When subject to high accelerating fields, copper cavities are limited to short-pulse applications. In contrast, SRF cavities have a much lower surface resistance, which implies low dissipation on the walls and high quality factors (of about $10^{10}$, compared to $10^4$ for copper) [13]. Taking into account the refrigerator power to keep the cavity in the superconductor state, SRF cavities are considerably more economical than copper cavities, and present huge benefits, especially for long-pulse applications. At high magnetic fields, however, high-temperature superconductor cavities can dissipate as much power as copper due to the nucleation and motion of vortices.

At low enough temperature and applied magnetic field (which for now we assume to be constant in time), superconductors exhibit the Meissner effect: magnetic fields are expelled from the interior of the superconductor, exponentially decaying from the interface surface. Larger applied magnetic field can destroy this Meissner state in two ways, depending on the type of superconductor. In type-I superconductors, an abrupt phase transition takes place at the thermodynamic critical field $H_c$, above which the superconductor is in the normal state. In type-II superconductors, the situation is slightly more complicated. Magnetic flux penetration starts, via vortex nucleation, at a lower magnetic field $H_{c1} < H_c$. $H_{c1}$ is called the lower critical field. The transition to the normal phase takes place at the upper critical field $H_{c2}$ ($H_{c2} > H_c$). In the intermediate range, $H_{c1} < H < H_{c2}$, the system is in the vortex lattice state.

1.2. The superheating field

For these cavities during operation, the external magnetic field is parallel to the superconductor surface. In many applications, the threshold field for flux penetration onto the superconductor is not set by $H_c$ or $H_{c1}$ (for type-I and type-II superconductors, respectively); it is set by the metastability limit of the Meissner state, i.e. by the superheating field [14–26]. The Meissner state is metastable at $H_c < H < H_{sb}$ for type-I superconductors, and at $H_{c1} < H < H_{sb}$ for type-II superconductors. The onset of instability of the Meissner state is related to the vanishing of a surface energy barrier that prevents field penetration onto the superconductor even when $H > H_c$ or $H > H_{c1}$.

$^7$ At higher magnetic fields ($>H_{c2}$), surface superconductivity can persist up to a third critical field, $H_{c3}$. This critical field should not be mistaken for the superheating field, below which the system displays bulk superconductivity and field expulsion.
The metastable Meissner state is analogous to the state of superheated water (perhaps explaining the name ‘superheating field’). Liquid water in a glass can be superheated in a microwave to a temperature above the liquid–gas transition temperature, but still remain in the liquid state due to the surface tension barrier at the liquid–gas interface, causing small vapor bubbles to contract rather than grow. Surface tension in water is analogous, for instance, to the surface tension due to the energy barrier preventing vortex nucleation in type-II superconductors. Unlike the case of water, as we argue in section 1.4, thermal nucleation of vortices occurs at relatively long time scales, suggesting that the Meissner state can be sustained in RF applications for fields as large as the superheating field. However, this scenario can considerably change when one considers the effects of disorder in the superconductor. Section 4.3 discusses disorder-induced nucleation of vortices.

The superheating field is associated with spinodal curves where the local stability of the Meissner state is broken. This is a more precise definition that is useful for both type-I and type-II superconductors. We shall discuss calculations of the superheating field in section 2. Our calculations there will be assuming an external field that is constant in time and ignore thermal fluctuations. We here discuss these approximations.

1.3. Why GHz is slow

Calculations of the superheating field for DC applied magnetic fields will be accurate for RF applications when the microscopic relaxation times are smaller than the time scales that are associated with changes in the fields inside the cavity. Time scales for the latter are of order of nanoseconds [13]. A version of time dependent Ginzburg–Landau theory given by Gor’kov and Eliashberg predicts the characteristic relaxation time near $T_c$: $\tau_{GL} = \pi \hbar / [8 k (T_c - T)]$, where $\hbar$ is the Planck constant divided by $2 \pi$ and $k$ is the Boltzmann constant [5]. For $T_c - T = 1$ K, one obtains $\tau_{GL} \sim 10^{-3}$ ns for oscillating fields parallel to the vortex surfaces. Using $\Delta \sim k T_c$, where $\Delta$ is the superconductor gap, we find $\tau_{GL} \sim \Delta^{-1}$ at low temperatures, which is similar to the scaling of collective modes in unconventional superfluids (see e.g. section 23.5 of [27]). However, note that Gor’kov and Eliashberg theory is applicable to gapless superconductors, filled with magnetic impurities and sufficient pair-breaking strength. For superconductors with a clean gap, the relaxation time is expected to be larger than $\tau_{GL}$, and to scale with the inelastic phonon-scattering time $\tau_{Ph}$, which, near $T_c$ is of the order of $\sim 10^{-8}$ s in Al and $\sim 10^{-11}$ s for Pb [5], due to its larger critical temperature. Yoo et al measured an ultra fast electron–phonon relaxation time of 360 fs for niobium [28]. So, at GHz frequencies we may ignore the time dependence in studying the stability.

1.4. Why thermal fluctuations are small

One key question for our purposes is whether thermal fluctuations can help activate vortices over the surface barrier. Thermal fluctuations in most superconductors (apart from the high-$T_c$ cuprate superconductors) are very small. This is due to the same approximation that makes the BCS theory of superconductors so successful. BCS theory is a mean-field theory of interacting Cooper pairs, which becomes exact when each Cooper pair interacts with an infinite number of neighbors (thus seeing the mean behavior of the system). Each Cooper pair is of radius roughly the coherence length $\xi$, so BCS theory will be valid when the density of Cooper pairs times $\xi^3$ is large. Simple estimates show that there are about $10^9$ centers of Cooper pairs within the region occupied by each pair state; a scenario where the pairs strongly overlap in space, and each pair only feels the average occupancy of the other pair states [35]. Thermal fluctuations of vortices will be unimportant so long as the condensation energy density—the amount of energy $F$ that is necessary to destroy superconductivity over a unit volume—times $\xi^3$, is large compared to $k_B T$. Table 1 gives the characteristic temperature $T_{th} = F\xi^3 / k_B$ where fluctuations will become important, for niobium and also three candidate materials being explored for next generation accelerating cavities. Only for NbN this characteristic temperature remotely comparable to $T_c$.

We can gain further insight from an analytic calculation of $E_c / (k_B T)$, where $E_c$ is the energy per unit length of a vortex line integrated over a coherence length $\xi$. Using results from BCS theory, the zero-temperature thermodynamical

Table 1. Representative material parameters for niobium, the traditional superconducting material used in SRF cavities, as well as candidate SRF materials that have the potential to reduce cooling costs due to their higher $T_c$. The coherence length $\xi$ is calculated using equations in [29]. The penetration depth $\lambda$ is calculated from equation 3.131 in [5]. The ratio $\kappa = \lambda / \xi$ is called the Ginzburg–Landau parameter, and determines many properties of superconductors. A residual resistivity ratio of 100 was assumed for niobium. For MgB$_2$, the values of $\lambda$ and $\xi$ are experimental values given in the reference. For calculations, $H_c = \phi_0 / (\mu_0 (2 \sqrt{2} \pi \xi \lambda))$ is used [5], $H_{1s}$ for Nb is found from fit to numerically computed data in [30, 31]. $H_{2s}$ for strongly type II materials is found from equation 5.18 in [5]. $H_{0s}$ is calculated using $H_{0s} \sim H_c (0.75 + 0.54 \xi^{-1/2})$ [14]. The condensation energy density $F$ is given by $\mu_0 H_c^2 / 2$ [5]. Nb data is extracted from [32], Nb$_3$Sn data from [30], NbN data from [33], and MgB$_2$ data from [34].

<table>
<thead>
<tr>
<th>Material</th>
<th>$\lambda$(nm)</th>
<th>$\xi$(nm)</th>
<th>$\kappa$</th>
<th>$T_c$(K)</th>
<th>$H_{1s}$(T)</th>
<th>$H_{c2}$(T)</th>
<th>$H_{0s}$(T)</th>
<th>$F$(J m$^{-3}$)</th>
<th>$F\xi^3 / k_B$(K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nb</td>
<td>40</td>
<td>27</td>
<td>1.5</td>
<td>9</td>
<td>0.13</td>
<td>0.21</td>
<td>0.25</td>
<td>17.547</td>
<td>25.009.0</td>
</tr>
<tr>
<td>Nb$_3$Sn</td>
<td>111</td>
<td>4.2</td>
<td>26.4</td>
<td>18</td>
<td>0.042</td>
<td>0.5</td>
<td>0.42</td>
<td>99.472</td>
<td>533.6</td>
</tr>
<tr>
<td>NbN</td>
<td>375</td>
<td>2.9</td>
<td>129.3</td>
<td>16</td>
<td>0.006</td>
<td>0.21</td>
<td>0.17</td>
<td>17.547</td>
<td>31.0</td>
</tr>
<tr>
<td>MgB$_2$</td>
<td>185</td>
<td>4.9</td>
<td>37.8</td>
<td>40</td>
<td>0.017</td>
<td>0.26</td>
<td>0.21</td>
<td>26.897</td>
<td>229.1</td>
</tr>
</tbody>
</table>

8 A simple estimate given in section 10.3 of [5], assuming a Debye phonon spectrum and free-electron Fermi surface, gives $\tau_{Ph}$ scaling as $T_c^{-3}$. 

Topical Review

critical field is given by $H_c(0) = 2 \sqrt{\pi} \sqrt{\mathcal{N}(0) \Delta}$, where $\mathcal{N}(0) = m k_F/(2 \pi^2 \hbar^2)$ is the density of states at the Fermi energy, $\Delta$ is the superconductor gap at zero temperature, and $k_F$ is the Fermi wave number. Also, $\Delta \approx 1.76 k_B T_c$ and the coherence length $\xi_0 = h v_F/(\pi \Delta)$, where $v_F$ is the Fermi velocity. Thus,

$$E_v \sim \frac{H_c^2 \xi_0^2}{k_B T} \approx \frac{1.4}{t} \left( \frac{\xi_0}{\Delta} \right)^2,$$

where $\xi_0 = \hbar v_F/(2 m)$ is the Fermi energy, and $t = T/T_c$. Since the gap is much smaller than the Fermi energy, we can neglect thermal nucleation of vortices; unlike the case of superheated water, the effects of thermal fluctuations is very small. More generally, we expect that $\tau_{\text{mic}} \ll \tau_{\text{cav}} \ll \tau_{\text{n.v.}}$, where $\tau_{\text{mic}}$, $\tau_{\text{cav}}$, and $\tau_{\text{n.v.}}$ correspond to time scales associated with microscopic degrees of freedom, the variation of the cavity fields, and thermal nucleation of vortices, respectively.

The negligible effects of thermal fluctuations tells us that estimating the limiting superheating field of a perfectly clean surface will not be analogous to bubble formation for superheated water. Instead, we shall use linear stability theory in section 2.2 to estimate the field at which the uniform Meissner state becomes energetically unstable to an infinitesimal perturbation in the space of magnetic fields and superconducting order. A variant of critical droplet theory will appear in section 4.3, where we estimate the effects of flaws and disorder in nucleating vortex penetration.

2. Basic theory of the superheating field

The superheating field $H_{\text{sh}}$ is set by the competition between magnetic pressure (imposed by the external magnetic field), the energy cost to destroy superconductivity, and the attractive force due to the zero-current boundary condition at the interface. In Ginzburg–Landau theory, the ratio $\kappa = \xi / \lambda$ of the penetration depth $\lambda$ to the coherence length $\xi$ determines many properties of superconductors. In particular, $\kappa < 1/\sqrt{2}$ and $\kappa > 1/\sqrt{2}$ are associated with type-I and type-II superconductivity, respectively. In the flux-line lattice of type-II superconductors, both the vortex supercurrent and magnetic field are confined to a tube of radius $\lambda$. The superconducting state is destroyed (the density of superconducting electrons vanishes) over a smaller vortex core of radius $\xi$. Within GL theory, $H_{\text{sh}}(T)/H_c(T)$ depends on materials properties only through the parameter $\kappa$, which is independent of temperature. A careful calculation using linear stability analysis [14] shows that $H_{\text{sh}}$ plateaus at about $0.75 H_c$ in the large $\kappa$ limit, and diverges as $\kappa^{-1/2}$ for $\kappa \ll 1$.

2.1. Simple arguments for the superheating field

We now give simple arguments and pictures to estimate the superheating field of superconductors (see e.g. [36]). The main idea is to compute the work necessary to push magnetic field onto the superconductor through an energy barrier set by the magnetic energy, and compare the result with the condensation energy. It is worth noting that there are important qualitative differences between these simple arguments and the actual linear stability analysis of the GL free energy. We will return to these issues when we discuss the effects of anisotropy in section 4.2, and discuss them further in the full publication [36].

Consider a superconductor occupying the half-space $x > 0$, and subject to an applied magnetic field $H$ that is parallel to its surface, along the direction $z$. We illustrate this geometry on the left side of figure 1, where ‘SC’ stands for superconductor. Note that the superconductor region extends to infinity in the positive and negative $y$ and $z$ directions, and in the positive $x$ direction; there are no ‘corners’ in this geometry.

Let us start with the argument for the superheating field of a type-I superconductor. For small external magnetic fields, the order parameter does not vanish at the vacuum-superconductor interface. However, if we push a slab of magnetic field onto the superconductor (just enough to make the order parameter vanish at the interface), we will destroy superconductivity over a length scale of order $\xi$. The work per unit area that is necessary to push magnetic energy onto the superconductor is set by the magnetic pressure and the penetration length; it is given approximately by $[H_{\text{sh}}/(4\pi)] H_{\text{sh}} \lambda$ in cgs units. To estimate the superheating field, we compare this work with the condensation energy per unit area $[H_c^2/(8\pi)] \xi$, resulting:

$$\frac{H_{\text{sh}}}{H_c} \approx 2^{-1/2} \kappa^{-1/2}. \quad (2)$$

Equation (2) should be compared with the small-$\kappa$ limit of the exact result using Ginzburg–Landau theory [14]: $H_{\text{sh}}/H_c \approx 2^{-1/4} \kappa^{-1/2}$.\footnote{The absence of corners is an important limiting factor in our approach, for corners typically facilitate field penetration in real samples of arbitrary shapes. Modern RF cavities have an approximate cylindrical shape in the region of high magnetic fields (see right side of figure 1), with no corners, so such geometric considerations become unimportant.}

\[ \text{Figure 1. (On the left) Illustration of a superconductor occupying the half-space } x > 0, \text{ and subject to an applied magnetic field } H \text{ that is parallel to the } z \text{ axis. ‘SC’ stands for superconductor. (On the right) Approximate shape of a superconducting RF cavity in the regions of high magnetic fields. As in the flat case, the magnetic field that is generated by the accelerating beam (and excited by an external RF source, driving the operating/accelerating mode) is parallel to the interior surface of the cavity.} \]
In type-II superconductors, field penetration occurs via vortex nucleation, and the superheating field is set by the magnetic pressure that is necessary to push a vortex through a surface barrier onto the superconductor. There are two steps to this penetration. First, the core of the superconducting vortex (of radius $\sim \xi$) must penetrate into the surface, at a cost given by the core volume times the condensation energy. Second, this newly penetrated vortex must fight past an attractive force toward the surface due to the boundary conditions at the surface, which is usually estimated by the attraction to an ‘image vortex’. Below we discuss the superheating field estimated from the initial penetration of the vortex. (Bean and Livingston’s original estimate [26] of the superheating field starts (somewhat arbitrarily) at a distance $x = \xi$ after this initial penetration, and focuses on the effects of the attractive longer-range force.)

Figure 2 illustrates the penetration of a vortex core (red disk) onto a superconductor occupying the half-space $x > 0$. The magnetic work per unit length to push the vortex core onto the superconductor is given approximately by the condensation energy (per unit length):

$$\frac{H_{bh} \Phi_0}{4 \pi \pi^2} \approx \frac{H_c^2}{8 \pi \pi^2},$$  \hspace{1cm} (3)

where $H_{bh}/(4 \pi)$ is the magnetic pressure, $\Phi_0$ is the fluxoid quantum, $\pi^2$ is the vortex area in the $xy$ plane, $4 \pi \xi$ is approximately the area that is associated with the region of field penetration (area of the orange box in figure 2); it is the amount of the area of the vortex that penetrates the superconductor when a vortex core is pushed inside, and $\pi^2$ is the area of the vortex core. Using $\Phi_0 = 2 \sqrt{2} \pi H_c \xi$ in equation (3):

$$\frac{H_{bh}}{H_c} \approx \frac{\sqrt{2}}{32} \approx 0.14,$$  \hspace{1cm} (4)

independent of $\kappa$.

How does this estimate compare with the field estimated from the attractive force, and with the true answer? The true answer, given below in section 2.2, is about five times larger: $H_{bh}/H_c \approx 0.75$. Bean and Livingston’s estimate of the superheating field due to the attractive force to the image vortex is $H_{bh}/H_c = 0.71$, of the same form as our estimate 0.14 but larger and closer to the true estimate. We present the calculation of the field necessary to introduce the core primarily due to its simplicity, and also because it motivates our analysis of anisotropic superconductors in section 4.2.

One should think of these two contributions as being sequential rather than serial: first the core must penetrate, and then the vortex must fight the longer-range attraction to enter the bulk. (It is interesting and convenient that these two fields are of the same scale.) The GL calculation in section 2.2 of course incorporates both the initial core penetration and the longer range attractive force, together with cooperative effects of multiple vortices entering at the same time.

Note that, while the field needed to push the vortex core into the superconductor is roughly comparable to that needed to push the vortex past the attractive long-range potential, the two contributions contribute very differently to the total energy barrier to flux penetration. Energy is force times distance: the two forces are comparable but the Bean–Livingston force acts on a scale longer by a factor $\kappa = \lambda/\xi$ than our core nucleation, and will dominate the barrier height. Finally, that in practice the dominant mechanisms for vortex nucleation that set the superheating field will not involve straight vortices penetrating all along their lengths (as in our calculation above) or, even more impressively, arrays of straight vortices cooperatively pushing their way through the surface barrier (section 2.2 below). We expect that disorder and flaws (discussed in section 4.3) will lead to localized intrusions of single vortex loops into the material (figure 8).

2.2. Linear stability calculation of the superheating field

In this section we have seen that the superheating field arises in a bulk superconductor due to the competing effects of magnetic pressure and the destruction of superconductivity. Using relatively simple arguments, we derived the qualitative dependence of this field on $\kappa$. We now describe a more rigorous calculation of the superheating field using a linear stability analysis. Linear stability analysis is commonly used in a variety of pattern formation problems [39–44]. For type II superconductors, the transition from the Meissner state to the mixed state is triggered by fluctuations of a critical wavelength that spontaneously break the transverse symmetry of the bulk sample, which when coupled to the inhomogeneous depth dependence of the Meissner state, make the
superheating transition a challenging application of this method. We here describe this calculation using the Ginzburg–Landau theory for concreteness, although the basic procedure could be extended to other theories as we discuss below. Our presentation follows closely the procedure described in [14], however, the calculation has a long history in the literature[18–25].

$$\delta^2 F = \int_0^{\infty} dx \left( \delta f \delta q_x, \delta f \delta q_y \right) \left( \begin{array}{c} -\xi^2 \frac{d^2}{dx^2} + q^2 + 3f^2 + \xi^2 k^2 - 1 \\ 2fq \\ 0 \end{array} \right)$$

where $k$ is the wave-number of the Fourier mode. The remaining Fourier components (corresponding to replacing $\cos \rightarrow \sin$ and vice-versa in equation (8)) are redundant as they decouple from those given in equation (8) and satisfy the same differential equations derived below.

After substituting into the expression (7) for the second variation and integrating by parts, we arrive at

$$\delta^2 F = \int_0^{\infty} dx \left( \delta f \delta q_x, \delta f \delta q_y \right) \left( \begin{array}{c} -\xi^2 \frac{d^2}{dx^2} + q^2 + 3f^2 + \xi^2 k^2 - 1 \\ 2fq \\ 0 \end{array} \right)$$

This matrix operator is self-adjoint, and the second variation will be positive definite if its eigenvalues are all positive. In the eigenvalue equations for this operator, the function $\delta q_y$ can be solved for algebraically. The resulting differential equations for $\delta f$ and $\delta q_y$ are

$$-\xi^2 \delta f'' + (3f^2 + q^2 - 1 + \xi^2 k^2) \delta f + 2fq \delta q_y = E\delta f,$$

and

$$-\lambda^2 \frac{d}{dx} \left[ \frac{f^2 - E}{f^2 + \lambda^2 k^2 - E} \delta q_y \right] + f^2 \delta q_y + 2fq \delta \delta f = E\delta q_y,$$

where $E$ is the stability eigenvalue. Note that by decomposing in Fourier modes, the two-dimensional problem is transformed into a one-dimensional eigenvalue problem. Numerically, it can be solved by the same methods as the Ginzburg–Landau equations [14].

The stability eigenvalue will depend on the solution of the Ginzburg–Landau equations, i.e., the applied magnetic field $H_0$ and the Fourier mode $k$ under consideration. The superheating field is found by varying both the applied magnetic field and Fourier mode until the smallest eigenvalue first becomes negative. The wave-number of the destabilizing fluctuations are therefore found simultaneously with $H_0$ and denoted by $k_c$. Values of $H_0$ and $k_c$ were calculated in Ginzburg–Landau theory for a wide range of $\kappa$ in [14, 25] along with analytic estimates. The results are summarized in figure 3.

For small $\kappa$, the critical fluctuation occurs with wave-number $k_c = 0$ while for large $\kappa$, $k_c > 0$. Interestingly, the transition to nonzero $k_c$ occurs at some critical $\kappa_c$ that is distinct from the type-I/type-II boundary ($\kappa = 1/\sqrt{2}$). Estimates of $\kappa_c$ vary in the literature from 0.5 [18] to 1.13(±0.05) [23]. Estimates of $k_c$ from solving equations (10) and (11) range from 1.10 [22] to 1.1495 [14] (our high-accuracy result).

The linear stability approach described in this section could be extended to other geometries as was done for the case of a superconducting film separated from a bulk superconductor by a thin insulating layer in [45]. More complicated theories of superconductivity can also be solved using our
methods by replacing the Ginzburg–Landau free energy with the appropriate analog, such as the Eilenberger formalism described in more detail in section 4.1.

3. Experiments

3.1. High power pulsed RF experiments

Some of the earliest measurements showing $H_{sh} > H_c$ for niobium were reported by Renard and Rocher based on DC magnetization measurements. Yogi et al performed a more systematic study at RF frequencies on samples of Sn, In, Pb, and alloys, in order to cover a range of $\kappa$ values [37]. Analysis of their data resulted in the vortex line nucleation model discussed in footnote 9. Noting that measurements of the RF critical field have shown inconsistency, Campisi used a very high power RF source at SLAC to very quickly ramp up the fields in cavities [46]. The goal of these high power RF measurements is to reduce the influence of defects by out-pacing the thermal effects they cause. Campisi performed high power RF measurements on Nb, Nb3Sn, and Pb cavities. Hays and Padamsee performed similar measurements on these materials at Cornell [47]. The niobium results are reproduced in figure 4, showing fairly reasonable agreement with the expected superheating field close to $T_c$ [11], but then diverging at lower temperatures.

After these experiments were performed, new preparation techniques were developed for niobium cavities, including a recipe involving electropolishing and a bake at 120 C. This recipe was found to avoid the ‘high field Q-slope’ (HFQS) degradation mechanism that occurs in niobium cavities at peak fields of approximately 100 mT [48, 49]. Experiments by Valles show that pulsed measurements of unbaked niobium produced curves that diverged from the expected $H_{sh}$ near the expected onset field of HFQS. However, after the bake was performed, the data agreed very well [50]. The $H_{fi}$ and $H_{sh}$ curves plotted in the figure were calculated from niobium material parameters that were extracted from measurements of $R_f$ versus $T$ and $f$ versus $T$ via the SRIMP Matthis–Bardeen code [51, 52]. The baked curve has a lower $H_{sh}$ due to the change in the mean free path after the bake, which in turn affects $\kappa$.

3.2. DC flux penetration measurements by N. Valles

Valles also performed measurements of the superheating field of unbaked niobium using a DC probe to avoid the effects of HFQS. Using a superconducting solenoid, he applied a DC

\footnote{11 $T_c$ assumed to be 9.2 K for Valles’ data.}
field to the exterior of a niobium cavity operating at low fields. A sudden decrease in the quality factor of the cavity indicated that flux from the magnet had penetrated to the interior cavity surface. The penetration field extracted from measurements of the applied field agreed well with the expected superheating field for unbaked niobium, as shown in figure 4 [50].

4. Beyond Ginzburg–Landau: Eilenberger, anisotropy, and disorder

The isotropic Ginzburg–Landau analysis of section 2 is a trustworthy estimate for the superheating field only for ideal surfaces of single-band superconductors with cubic symmetry near the superconducting transition temperature $T_c$. In this section we pursue three topics that introduce new physics to this calculation. First, superconducting RF cavities are usually run at temperatures significantly lower than $T_c$; niobium cavities, with $T_c \sim 9$ K are usually run at $T = 2–4$ K in working accelerators. In section (4.1) we review calculations of the superheating fields that use Eilenberger theory, which is valid at lower temperatures, presenting the analytic results [15] at large $\kappa$. Our estimates suggest that these Eilenberger corrections to GL are quantitatively important at operating temperatures, but not large. Second, many of the potential new superconductors have rather anisotropic crystal structures and electronic properties; if the superheating field has significant anisotropy, this could motivate single-crystal or controlled growth conditions to control surface orientations in cavities. In section 4.2 we review calculations [36] which show that this anisotropy will be small near $T_c$; we also discuss conflicting results for the anisotropy of multi-band superconductors (like MgB$_2$) at low temperatures. Third, in section 4.3 we estimate the effects of disorder and flaws in these materials, presenting both qualitative and simple quantitative estimates of the effects of defects and dirt in locally lowering the barrier to magnetic flux penetration and thus lowering the effective superheating field.

4.1. Eilenberger theory for lower temperatures

The Ginzburg–Landau approach to superconductivity is generally accurate near the critical temperature $T_c$, but usually the accuracy of its prediction worsen as temperature is lowered below $T_c$. A basic example of its failure is given by the temperature dependence of the order parameter $\Delta$: according to GL theory, $\Delta(T)$ behaves as $\sqrt{1 - T/T_c}$, in agreement near $T_c$ with the microscopic BCS theory. The latter, however, predicts that at low temperatures the order parameter is temperature-independent up to exponentially small corrections. For our purposes, the limited validity of GL theory implies that the dependence of the superheating field on $\kappa$ discussed in section 2 cannot be assumed to be quantitatively accurate at the low temperatures at which RF cavities are usually operated. This motivates us to consider a more general approach, valid at arbitrary temperature.

For low-$T_c$ superconductors, the coherence length $\xi_0 = \hbar v_F / 2\Delta_0$ is much longer than the Fermi wavelength; here $\xi_0$ is the zero-temperature coherence length for a clean superconductor with zero-temperature order parameter $\Delta_0$ and Fermi velocity $v_F$. Thanks to the separation in length scales (or equivalently, the separation in energy scales between $\Delta_0$ and the much larger Fermi energy), these superconductors can be modeled using the so-called quasiclassical approach, reviewed for example in [53, 54]. This powerful approach is quite flexible, permitting in principle to include effects such as Fermi surface anisotropy and impurity scattering (we will comment on the latter at the end of this section). This come at the price of having to calculate various Green’s functions from which physical quantities such as the order parameter and the current can be obtained. Such calculations are usually much more involved than that of the GL approach.

It was shown by Eilenberger [6] that one can arrive at an expression for the thermodynamic potential as functional of order parameter $\Delta(r)$ and vector potential $A(r)$, similar to the GL functional, once the quasiclassical equations for the Green’s function have been solved. While a general solution is not possible, for the case of a clean superconductor with spherical Fermi surface we developed in [15] a perturbative approach valid for large $\kappa$. Then the thermodynamic potential $\Omega$ is

$$
\Omega = \nu \int d^3r \left\{ \frac{1}{3} (\nabla \times A - H_0)^2 + \Delta^2 \log \left( \frac{T}{T_c} \right) \right. \\
+ \int (dn) \left[ \frac{\Delta^2}{\omega_n} - 2(\sqrt{\Omega_n^2 + \Delta^2} - \omega_n) \right. \\
+ \left. \frac{1}{\kappa_0^2} \Delta^2 \frac{\sqrt{\Omega_n^2 + \Delta^2}}{4\Omega_n^2} (n \cdot \nabla \phi_s^{(0)})^2 \right\}. 
$$

(12)

In this expression $\nu$ is the density of states at the Fermi energy, lengths are in units of the zero-temperature penetration depth $\lambda_0$,

$$
\frac{1}{\lambda_0} = \frac{8\pi}{3} \left( \frac{2\pi \xi_0}{\Phi_0} \right)^2 \nu \Delta_0^2.
$$

(13)
the vector potential is in units of $\Phi_0/2\pi \xi_0$ with $\Phi_0$ the magnetic flux quantum, $\kappa_0 = \lambda_0/\xi_0$, and $n$ is the unit vector on the Fermi surface. We also use the short-hand notations

$$\int (dn) = 2\pi T \sum_n \int \frac{dn}{4\pi}, \quad \Omega_n = \omega_n - in \cdot A. \tag{14}$$

with $\omega_n = 2\pi T (n + 1/2)$, $n = 0, 1, 2, \ldots$, the fermionic Matsubara frequencies, and

$$s^{(0)} = \frac{2\Delta}{\sqrt{\Omega_n^2 + \Delta^2}}. \tag{15}$$

The thermodynamic potential in equation (12) reduces to the GL one near $T_c$\textsuperscript{12} and it can be used to find the superheating field at arbitrary temperature in the regime $\kappa_0 \gg 1$. The calculation of $H_{ab}$ proceeds in the same manner as in the GL approach, by studying the stability against small perturbation of the local minima of $\Omega$. This study was performed in [15] at leading order in $\kappa_0 \rightarrow +\infty$. The ratio $H_{ab}/H_c$ between superheating and critical field can be calculated analytically at $T = T_c$ and $T = 0$:

$$\frac{H_{ab}^\infty}{H_c}(T_c) \simeq 0.745, \quad \frac{H_{ab}^\infty}{H_c}(0) \simeq 0.840, \tag{16}$$

where we use the $\infty$ symbol in the superscript to indicate that these are leading-order results. Interestingly, the zero-temperature ratio is almost 13% larger in the regime $\kappa_0 \gg 1$, indicating that naive extrapolation to low-temperatures of the GL result underestimates the superheating field. At arbitrary temperature, the $H_{ab}^\infty/H_c$ ratio can be found numerically and is shown in figure 5. Note the non-monotonic dependence of $H_{ab}^\infty/H_c$ on temperature, which leads the superheating field to acquire its largest value $H_{ab}^\infty \simeq 0.843H_c(0)$ at $T \approx 0.04T_c$.

It should be noted that while the Meissner state remains metastable up to $H_{ab}$, a clean superconductor can become gapless at a lower field $H_g$ [55]; for example at $T = 0$ we have $H_g \approx 0.816H_c < H_{ab}$. The field $H_g$ is relevant to applications such as superconducting cavities because as the applied field approaches $H_g$, AC losses rapidly increase. Indeed, in the presence of the AC losses are in general exponentially suppressed, but this ‘protection’ from losses is absent in the gapless state.

The above results are restricted to the leading order in $1/\kappa_0$, which makes it possible to neglect the contributions from the last term in square brackets in equation (12). At next to leading order, that term must be taken into account and leads to an expression for the superheating field of the form\textsuperscript{13}

$$H_{ab}(T) \approx \frac{H_{ab}^\infty}{H_c}(T) + \frac{h(T)}{\sqrt{\kappa_0}}. \tag{17}$$

This formula, with a weakly temperature-dependent dimensionless coefficient $h(T)$, has the same inverse square root dependence on $\kappa_0$ as the GL expression [14].

\textsuperscript{12} In considering the limit $T \rightarrow T_c$ in [15], a prefactor was missed in equation (29) and consequently equation (31), which should read respectively: $\eta_{gK} = 2\pi T \frac{2\sqrt{3}}{\xi_0} \approx 1.50\kappa_0$ and $\zeta(T) = \sqrt{2/3} \Delta_0/\Delta(T)\xi_0$ in the notation of that work.

\textsuperscript{13} This formula can be obtained by extending to next-to-leading order the calculations of [15] (Catelani, unpublished).

Figure 6. Phase diagram of anisotropic superconductors in terms of mass anisotropy ($\gamma = \sqrt{m_c/m_a}$) and GL ($\lambda_0/\xi_0$) parameters. The superconductor is of type-I to the left of the blue line, of type-II to the right of the red line, and mixed in between (in the mixed phase, the SC is of type-I for $c || z$ and of type-II for $c \perp z$). The blue and yellow regions correspond to the asymptotic solutions $H_{ab}^\infty/H_c \approx \gamma^{1/2}$ and $H_{ab}^\infty/H_c \approx 1$, respectively (within 10% accuracy). Note that the superheating field of MgB$_2$ is nearly isotropic near $T = T_c$. Reprinted figure with permission from [36]. Copyright 2016 by the American Physical Society.

In closing this section, let us comment briefly on the effect of impurity scattering. Both non-magnetic and magnetic impurities were considered in [55] in the limit $\kappa \rightarrow \infty$. At sufficient strength of the non-magnetic impurities scattering rate, there are some qualitative changes: the non-monotonicity of $H_{ab}(T)$ is suppressed, and more importantly the gap remains open up to $H_{ab}$. However, quantitatively the value of $H_{ab}/H_c$ is changed by at most a few percent. In contrast, adding magnetic impurities strongly decreases $H_{ab}$ similar to the well-known suppression of $T_c$ due to the pair-breaking effect of such impurities.

4.2. Anisotropic superconductors

Layered superconductors can display highly anisotropic critical fields. The upper-critical field of BSCCO\textsuperscript{14}, for instance, can vary by two orders of magnitude depending on the angle between the crystal anisotropy axis $c$ and the applied magnetic field [5]. Near zero temperature, the upper critical field of magnesium diboride is about six times larger for $c \perp B$ than for $c || B$ (see e.g. [56, 57]). Here we review [36], which investigates the effects of crystal anisotropy on the superheating field of superconductors, motivated partly by the opportunity of controlling surface orientation in order to achieve higher accelerating fields inside the cavity.

Near the critical temperature, for the anisotropy axis $c$ aligned with one of the Cartesian directions, the anisotropic formulation of Ginzburg–Landau theory [58–61] provides a clean approach to study the anisotropy of the superheating fields...
field. We can use a change of coordinates and rescaling of the vector potential to turn the anisotropic GL free energy onto isotropic form, and then use previous results from [14] to calculate the superheating field anisotropy of several materials. We find that:

$$H_{sh}^{\text{an}} = \begin{cases} H_{sh}^c(\kappa), & \text{for } c \parallel z, \\ H_{sh}^c(\kappa\gamma), & \text{for } c \parallel x \text{ or } y, \end{cases}$$

where the superheating field on the right-hand side is the solution of the linear stability analysis for isotropic Fermi surfaces, which we discussed in section (2.2), using $\kappa = \kappa_i$ and $\kappa = \kappa_c = \gamma\kappa_i$ for $c$ parallel and perpendicular to $z$, respectively. Within GL theory, $\gamma = \sqrt{m_c/m_s} = \lambda_c/\lambda_s = \xi_y/\xi_z$, with $m_c$, $\lambda_i$ and $\xi_i$ representing the effective mass, penetration depth, and coherence length along the $i$th direction, respectively. Since $H_{sh} \approx 0.75 H_c$ goes to a constant for large $\kappa$, we find that the superheating field is nearly isotropic for most high-$\kappa$ unconventional superconductors. On the other hand, $H_{sh} \approx 0.84 H_c\kappa^{-1/2}$ for small-$\kappa$ type-I superconductors, resulting in an anisotropy of about $\gamma^{1/2}$ when $\kappa_i\gamma$ is small.

Figure 6 displays a phase diagram in terms of $\kappa_i$ and $\gamma_i$, showing the region where GL theory predicts type-I (left of the blue line), type-II (right of dark red line) and mixed (in between dark red and blue lines) superconductivity, and the regions where each asymptotic solution is expected. Note, in particular, that $H_{sh}^{\text{an}}/H_{sh}^c \approx 1$ for MgB$_2$. This result is valid only near $T_c$, where the anisotropies in $\lambda$ and $\xi$ are equivalent. In the next paragraph we will use results from a two-gap BCS theory to estimate the superheating field anisotropy of MgB$_2$ at lower temperatures.

Theoretical and experimental studies indicate that the assumption $\lambda_c/\lambda_s = \xi_y/\xi_z$ (vortex and vortex core have identical shapes within GL theory) is strongly violated for low-temperature MgB$_2$, thus suggesting the use of two parameters to describe crystal anisotropy, namely $\gamma_c = \lambda_c/\lambda_s$ and $\gamma_c = \xi_y/\xi_z$. Also, $\gamma_c$ and $\gamma_c$ exhibit different temperature dependences, with $\gamma_c$ decreasing and $\gamma_c$ increasing for decreasing temperature, respectively. Calculations from [56] using a two-gap BCS model suggest that $\gamma_c$ and $\gamma_c$ become equal only at $T_c$, near zero temperature, $\gamma_c \approx 6$ whereas $\gamma_c \approx 1$, agreeing with some [57, 62, 63], but not all (see [56] and references therein) experimental estimates.

We can use our simple estimates of section 2.1 to make a qualitative prediction for the resulting anisotropy $H_{sh}^{\text{an}} / H_{sh}^c$ in the superheating field, when $\gamma_c \neq \gamma_c$ deviates from the single-band GL prediction. Now the anisotropic shape of the vortex and vortex core plays an important role (see figure 7(a)). When $c$ is in the $xy$ plane, as in figure 7(b), for instance, the superheating field is estimated from the work performed to push the black-dashed ‘box’ into the superconductor, which can considerably vary from $c||y$ (left) to $c||x$ (right). This leads to an estimate $H_{sh}^{\text{an}} / H_{sh}^c \approx \gamma_c / \gamma_c$. A second estimate generalizes the Bean and Livingston argument of the longer-vortex impurity to incorporate anisotropy, and leads to a slightly different result: $H_{sh}^{\text{an}} / H_{sh}^c \approx \gamma_c / \gamma_c$. Yet a third calculation, which we term ‘Extended GL’, yields an almost isotropic result, and is based on a direct linear stability analysis of the anisotropic GL free energy (see equation (7) of [36]) assuming unconstrained $\lambda$’s and $\xi$’s.

Table 2 summarizes our estimates of $H_{sh}$ for the three geometries, using experimental values for $H_c$ and $\kappa$ for MgB$_2$. Note that we correct numerical discrepancies of our first estimates in the second row of the table: ‘1st (corrected)’. The last column shows the maximum superheating field anisotropy according to each method. Most of the values of $H_{sh}$ are

![Figure 7](image-url)
as low as $H_{sb} \approx 0.24\text{T}$ for Nb [13]. We discuss the origin of these disparate predictions further in [36].

Our GL arguments for the superheating field anisotropy can be trusted near $T_c$: at large $\kappa$ the superheating field anisotropy is not a reason to control surface orientation. Our arguments at lower temperature and for multi-band superconductors are more speculative. The vortex core shape will surely change for $x \sim \xi$ due to the boundary conditions at the surface; the anisotropy in the long-range attraction in multi-band materials may be different from that of a simple anisotropic GL approach. It will be important to apply linear stability analysis to more sophisticated theories, such as multi-gap BCS or strong-coupling Eliashberg theory, especially in the face of the conflicting results shown in table 2.

4.3. Disorder and vortex nucleation

Niobium RF cavities are routinely operated in the metastable regime, at fields $H_{c1} < H < H_{sb}$ above the field $H_{c1}$ where vortices in equilibrium would penetrate into the superconductor (and dissipate roughly the same energy as in a normal metal). Table 1 in section 1.4 gives $H_{c1}$ and $H_{sb}$ for other candidate materials. For niobium this metastable regime gives us an important factor of $\sim 1.6$ in field. Running in the metastable regime is crucial for utility with the higher temperature superconductors, whose $H_{c1}$ equilibrium fields are much lower than the operating fields for current Nb cavities (table 1).

It took many years of experimentation to raise operating fields of the niobium cavities to approach near to their fundamental limits. Will the new, more complex materials be fundamentally more challenging to optimize? Our preliminary experimental cavities using Nb$_3$Sn appear already to be operating above $H_{c1}$ [64], but are not yet delivering anywhere near to the theoretically predicted superheating field. Just as we have been exploring the fundamental theoretical limits to the fields for ideal surfaces, in this section we explore the fundamental theoretical challenges in minimizing the effects of dirt, flaws, and defects in lowering the barriers to vortex entry.

What kind of flaw or disorder fluctuation would be needed to allow vortices to enter at fields substantially lower than the superheating field? How big a damage region is needed to bypass the surface barrier to vortex entry? Damage will significantly affect the superconducting properties if the flaw or fluctuation has a characteristic length of order the coherence length $\xi$. Since the proposed candidate materials for next generation SRF cavities have shorter coherence lengths than niobium (table 1), this potentially could imply that these new materials are more susceptible to defects and dirt.

Figure 8 shows a cartoon of a vortex loop entering a superconductor. Based on the discussion at the end of section 2.1 and the caption of figure 8, at external fields far from $H_{c1}$ and $H_{sb}$, the energy of the vortex loop will grow in the absence of disorder until it reaches a critical radius $R_c$, at which point the energy will again decrease. This critical radius and the needed damage zone will get smaller as the field $H$ grows, vanishing at $H = H_{sb}$.

The energy per unit length of the vortex loop will have two contributions—a curvature energy and an attractive energy between the vortex and the surface. The latter can be estimated from the attraction of a straight vortex to the ‘image vortex’ needed to set the correct boundary conditions at the surface. This potential barrier (the major component of the superheating field) was estimated by Bean and Livingston for high $\kappa$ type-II superconductors [26]. The unitless Gibbs free energy per unit length $4\pi G/(\sqrt{2} \ H_c \ \Phi_0)$ of a straight vortex flux line a depth $x$ inside a superconductor with external field $H$ can be written in the (London) large-$\kappa$ limit as [12]:

$$ G = \Phi_0 \left( H (e^{-x/\lambda} - 1) - \frac{\Phi_0}{2 \pi \lambda^2} K_0(2x/\lambda) + H_{c1} \right). $$ (19)

$$ \frac{4 \pi G}{\sqrt{2} \ H_c \ \Phi_0} = g(x) = h(e^{-x/\lambda} - 1) - \frac{K_0(2x/\lambda)}{2 \kappa} + \frac{\ln \kappa}{2 \kappa}, $$ (20)
where \( h = H / (\sqrt{2} H_c) \), and \( K_c \) denotes the modified Bessel function of imaginary argument [65], and for now \( \kappa \) and \( \lambda \) are the Ginzburg–Landau parameter and penetration length of the pure material, respectively. The first term is a magnetic pressure, the second term is the interaction with the ‘image vortex’ that imposes the correct boundary conditions, and the third term is the energy per unit length of a vortex deep in the superconductor. We can estimate \( R_c(H) \) by setting the derivative \( dG / dx = 0 \) in equation (19) and expanding the Bessel function for small arguments, leading to \( R_c(H) \sim \xi H_{sh}/H \). At the lowest field for vortex penetration \( H_{c1} \) this expansion is unreliable; however, since \( H_{c1} = H_{sh} \log(\kappa) / \kappa \) [12], the resulting estimate \( R_c(H_{c1}) \approx \xi / \log(\kappa) = \lambda / \log(\kappa) \) is still quite good, as shown in figure 9(a). But the new materials of interest have lower critical fields \( H_{c1} \) too small to be useful; we must run at fields \( H \) comparable to \( H_{sh} \). Near \( H_{sh} , R_c \approx \xi \), as again shown in figure 9. For disorder or defects to remove this energy barrier, they will thus necessarily have to strongly affect a region of volume \( \sim \xi R_c^2 \sim \xi^3 \).

To make this more quantitative, one needs to identify and model the dominant mechanism for vortex nucleation. If the characteristic defect size is large compared to \( \xi \) (e.g., nucleation on grain boundaries or inclusions of competing phases), one must model and control these individual defects. Clean grain boundaries are usually atomistically sharp (much thinner than \( \xi \)) and hence do not significantly decrease the local superconducting properties; indeed, studies of hot spots in large grain niobium cavities show no correlation with grain boundaries [66], and using single crystals to avoid grain boundaries has not improved performance [67, 68]. But in more complex materials, grain boundaries could be more disordered, thicker, or contaminated by impurities, and a grain boundary or grain boundary intersection with the correct orientation with respect to the surface could provide a route to entry. The effect of surface roughness on Bean and Livingston’s surface barrier has been studied in [69]. Kubo has used the London model to investigate the effects of nano-scale surface topography on the superheating field [70]. Perhaps most dangerous could be inclusions of metallic or poorly superconducting second phases, or irregularities in the surface morphology.

If the characteristic defect size is small compared to \( \xi \), and if the defects are uncorrelated in position, then the fluctuations in regions of order \( \xi^3 \) can be quantitatively estimated to linear order using the central limit theorem. This leads to Gaussian random fluctuations in the superconducting properties. For example, for alloys and doped crystals there are natural concentration fluctuations that will locally change the superconducting transition temperature, coherence length, condensation energy, and other properties. This is the traditional theoretical framework for field-theoretic calculations of the effects of disorder.

Let us hypothesize a system where the critical temperature is decreased due to disorder. In the context of Ginzburg–Landau theory for a homogeneous system, a change in the critical temperature yields a change in the coefficient \( \alpha = \alpha(\chi) \) of \( \psi^2 \), where \( \psi \) is the superconductor order parameter [5]. The probability of a fluctuation in \( \alpha(\chi) \) away from its pure value \( \alpha_0 \) would be proportional to

\[
\Pi [\alpha(\chi)] \propto \exp \left( -\int (\alpha(\chi)/\alpha_0 - 1)^2/(2\sigma^2) d^2x \right), \quad (21)
\]

where \( \sigma \) is a material-dependent constant that encapsulates the likelihood that the dirt in the material will cause a given fractional change \( \alpha/\alpha_0 \) in the critical temperature. The constant \( \sigma \) will become larger either if there are bigger concentration fluctuations or if the material is particularly sensitive to dirt. In principle, we should now calculate the most probable three-dimensional profile \( \alpha(\chi) \) needed to flatten the energy barrier and allow vortices in at a lowered field \( H < H_{sh} \), and then use \( \Pi [\alpha(\chi)] \) in equation (21) to estimate
The vortex will see a depth-dependent thickness $d$ (section 5).

The probability per unit surface area $P(H/H_{th}) = \Pi\{\alpha(x)\}$ of vortex penetration.

Rather than doing this full variational calculation, we build on the Bean–Livingston model of equation (19). In GL theory, the characteristic lengths scale as $\lambda \sim \alpha^{-1}$ and $\xi \sim \alpha^{-1/2}$. Hence we distinguish $\lambda_0$, $\xi_0$ and $\kappa_0$ for the pure material from $\lambda(x) = \lambda_0/a$, $\xi(x) = \xi_0/\sqrt{a}$ and $\kappa(x) = \lambda(x)/\xi(x) = \kappa_0/\sqrt{a}$ for the damaged region, where $a = \alpha/\alpha_0$.

What is the minimum amount of dirt that is necessary to reduce the superheating field to a certain value? For instance, how much dirt would it take to reduce $H_{th}$ for Nb$_3$Sn (estimated at 0.42 T in table 1) to $H = 0.25$ T ($H_{th}$ for niobium), a factor $H/H_{th} \sim 0.6$? One would need enough dirt to ‘flatten’ the surface barrier between $\xi_0$ and $R_c(H) \approx \xi_0 H_{th}/H = 5 \xi_0/\beta$ along the $x$ direction (thus allowing for vortex penetration), as shown in the dashed line of figure 9(a). In general, we are interested in finding an $x$-dependent parameter $\alpha(x)$ that flattens the energy barrier from $x = \xi_0$ to $x = x_f$, where $x_f > \xi_0$, and is defined by $G(x_f) = G(\xi_0)$. The solution for $\alpha(x)$ is then found from the equation $G(x) = G(\xi_0)$ for $\xi_0 < x < x_f$, and $\alpha(x) = \alpha_0$ for $x > x_f$, where in the left and right-hand sides we use $\{\lambda(x), \xi(x)\}$ and $\{\lambda_0, \xi_0\}$ in equation (19), respectively.

Note that we are making a rough approximation here. The magnetic fields and supercurrents surrounding the vortex line will see a spatially varying critical temperature $\alpha(x)$ whenever it is far from the surface, and properly measuring its energy and thus the surface attraction should include the resulting shift in energy. The depths $x$ of importance to us are of order the coherence length $\xi$, and thus these long distance fields and currents are largely canceled by the image vortex a distance $2x$ away. The vortex will see a depth-dependent disorder, but its energy will be qualitatively well described by our model in the region $H \sim H_{th}$.

Figure 9 shows $\alpha/\alpha_0$ as a function of $x/\xi$ for several values of $H$ in the interval [$H_{th}, H_{th}$]. Apart from an overall constant given by the normalization of the Gaussian, the negative logarithm of the probability of this fluctuation as a function of the lowered entry field $H$ is

$$-\log(P(H/H_{th})) = \Pi\{\alpha(x)\}$$

$$= \int (\alpha(x)/\alpha_0 - 1)^2/(2\sigma^2) dx$$

$$= \frac{\xi_0^2}{2\sigma^2} \int (\alpha(\xi_0 u)/\alpha_0 - 1)^2 du$$

$$= \frac{\xi_0^2}{2\sigma^2} P(H/H_{th}),$$

a measure of the relative logarithmic reliability of the superconductor to disorder-induced nucleation. Here we pull out the volume $\xi_0^3$ of the damage zone by changing variables to $u = x/\xi_0$. In a three-dimensional system with a semicircular vortex nucleation approximation, we can use our Bean–Livingston style methods to approximate this as a one-dimensional integral

$$P_{3D}(H/H_{th}) = \int \pi u (\alpha(\xi_0 u)/\alpha_0 - 1)^2 du.$$  

(26)

For a vortex pancake nucleation event for a thin SIS film of thickness $d$ (discussed in section 5), we find

$$P_{2D}(H/H_{th}) = (d/\xi_0) \int (\alpha(\xi_0 u)/\alpha_0 - 1)^2 du$$

(27)

(see figure 10).

Clearly, the relative reliability decreases rapidly as $H$ approaches $H_{th}$, by many orders of magnitude in this model calculation. The high-$\kappa$ calculation of Bean and Livingston cannot be simply extrapolated to niobium, but there is no reason to doubt that a similar sensitivity of the barrier to disorder in superconductors is much stronger than its dependence on $\kappa$ or $\xi$. This suggests, examining figure 10, that the factor of five to ten change in $\xi_0$ with the new superconductors may not be so dangerous. The resulting two to three orders of magnitude smaller volume for the critical damage zone at fixed field, it would seem, could be remedied by working not at $0.8H_{th}$ but at perhaps $0.6H_{th}$ (figure 10).

Manufacturing high-quality cavities from these new materials may be challenging. What our calculation can provide is reassurance that these materials should not be avoided because of their shorter coherence lengths.

5. Laminates and vortex penetration

In recent years, much effort in superconducting RF has been devoted to exploring single or multiple thin films—laminated structures hopefully tunable to optimize performance. This section is devoted to exploring possible advantages to such
laminates. The work in this section relies heavily on extensive discussions and consultation with Alex Gurevich, whose work prompted most of the calculations presented.

In practical terms, two of the candidate materials (Nb$_3$Sn and NbN) can be grown by deposition on Nb surfaces, so fabricating a surface layer onto a Nb cavity leverages existing expertise. Gurevich points out [72] that thermal conductivities of new candidate materials are often small; since the heat generated by the surface residual resistance at the surface must be conducted through the cavity, keeping the thickness of these new materials small can improve performance. (For Nb$_3$Sn, recent surface resistances have been small enough, at least at low fields, that thickness may not be an issue.) Gurevich has also proposed [73] separating one or more superconducting layers by insulating layers (a SIS geometry). Calculations show [45] that laminates do not substantially improve the theoretical maximum superheating field in AC applications beyond that of pure materials (or thick layers) for the film-insulator-bulk structure,\footnote{A free-standing superconducting layer (or a layer surrounded by insulators) with thickness small compared to the magnetic penetration depth $\lambda$ can have an enormous superheating field (since it can remain superconducting without paying most of the cost of expelling the flux). In the accelerator community, there is widespread focus on raising this $H_{\text{c1}}$ for the superconducting film [74, 75]—defined, somewhat unphysically [76] as the minimum field needed for a vortex to be stable parallel to and inside the film. But such an in-film stable vortex configuration demands magnetic flux on both sides of the film. In a GHz AC application, pushing the flux through the film twice per cycle generates unacceptable heating [76]. Besides, any such parallel vortex would be precariously unstable to formation of two vortex pancakes. A thin superconducting layer with a large magnetic penetration depth atop a lower $H_{\text{c1}}$ layer with a small penetration depth can have modestly higher superheating fields, due to the way the bottom layer modifies the magnetic field penetration.} though adding a thin S$'$ layer on the bulk S superconductor may lead to an enhancement of the energy barrier [77, 78].

Gurevich has suggested that the SIS geometry may have a different advantage—reducing the impact of flux penetration. Our calculations in section 5.3 suggest that SIS films with thickness $d$ small compared to the London penetration depth $\lambda$ will be more susceptible to vortex penetration than bulk films; the damage zone needed for vortex nucleation at fields below pure $H_{\text{c1}}$ can be thinner by the fraction $\lambda/d$, presumably making them much more likely. Also, one would naively expect it to be harder to grow low-defect two-layer laminates than depositing a single layer or preparing a pure surface. Layers thick compared to the penetration depth would presumably behave similarly to a bulk material; vortices deeper than $\lambda$ do not ‘feel’ the surface except insofar as other vortices penetrating the surface push them deeper.

The dynamics after flux penetration will be substantially different for the SIS geometry than for a simple 3D superconducting surface. In either case, a flaw may nucleate one to several vortex entries when the field increases in one direction; some or all may be ‘pulled back’ as the field shifts to the opposite direction. If the nucleation center flaws are rare and the vortices do not build up over time, they need not cause local heating enough to cause a quench. But since the RF cavities operate at GHz frequencies (billions of cycles per second), and each flaw could (or should) generate multiple vortices per cycle, potentially billions of vortices per second could be introduced by a single flaw if they can escape away from the defect and avoid re-annihilation.

In three-dimensions, a vortex penetrating at a point $(y, z)$ on the surface will grow in the $z$ direction pointing along the field as it penetrates a depth of order $x \sim \lambda$ (figure 11 left). If multiple vortices enter, they may push and entangle one another; as they interact with disorder in the material they may exhibit avalanches [79, 80]. During the field reversal, the points where the vortices exit the material will be forced together along the $z$ direction (shrinking in length), and new vortices with opposite winding number will nucleate (potentially annihilating some or all of the old vortices). Even if this process is incomplete, leaving some tangle of vortex loops, it may enter a kind of limit cycle. Indeed, many periodically stressed disordered dynamical systems can enter into limit cycles at low levels of stress, with a transition to ‘turbulent’ aperiodic behavior at a critical threshold (colliding colloids in reversing low-Reynolds number flows [81], plasticity in vortex structures of superconductors [82–84], etc). It is possible that the quench of RF cavities explores precisely this

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**Figure 11.** Vortices in a bulk superconductor for semiloops (left). Vortices in thin superconducting films separated by insulators form pancakes.**

---

**Figure 12.** For initial separations smaller than the impact parameter $x_{\text{imp}}$, the pancakes annihilate (inner solid, dash, dot), but larger than $x_{\text{imp}}$ they can wander away (dash–dot, outer solid). $\alpha = 0.2$ shown.
kind of dynamical phase transition, separating a local hot spot from an invading front of vortices. Apart from these brief speculations, we will not discuss three-dimensional dislocation dynamics further in this work; the remainder of this section will focus on the SIS geometry.

In the two-dimensional SIS geometry, a vortex penetration event may end with the vortex trapped in the insulating layer, leaving two 2D vortices penetrating the outer superconducting film (figure 11 right. See also footnote 15). Such 2D vortices, called pancake vortices, have been studied in great detail [12] in the context of high temperature cuprate superconductors, some of which are well described as nearly decoupled 2D superconducting sheets. A vortex pair nucleated by a defect at \((y, z)\) on the surface will separate along the \(z\) direction as the field increases, be buffeted by thermal fluctuations, dirt, defects, and other vortices as they separate, and then be pulled back along the \(z\) direction as the field reverses. (Some of the other vortices will be emitted by the same defect, once the initial pair departs and the resulting long-range suppression of nucleation drops, see section 5.4.) In this part we shall explore Gurevich’s suggestion that, even after billions of cycles, this annihilation should be effective at avoiding vortex escape (presumably preventing a buildup of vortices which otherwise would lead to a quench).

In section 5.1, we introduce an ‘impact parameter,’ the amount of lateral vortex separation between a vortex–anti-vortex pair that can be tolerated during a cycle while still expecting them to annihilate, in section 5.2, we examine the expected lateral meandering distance expected from pancake vortices in an RF cycle, in section 5.3, we examine the expected meandering due to disorder, and in section 5.4, we briefly consider the effect of vortex–vortex interaction and the situation of two nearby defects.

5.1. Impact parameter

How far \(\Delta x\) perpendicular to the field must a vortex pair migrate before their mutual attraction ceases to be strong enough to annihilate them at the end of a cycle? Figure 12 shows the trajectories for a pancake pair as they return at the end of a cycle, separated by different distances \(x\) perpendicular to the external magnetic field, using the vortex interaction formulation from [85]. There is a separatrix between trajectories which collide and trajectories which miss each other. We will call the value of \(\Delta x\) at this separatrix the impact parameter, \(x_{\text{imp}}\). For perhaps credible parameters \(d = 30\,\text{nm}, \lambda = 100\,\text{nm}, \mu B H_{\text{sh}} = 0.4\,\text{T}, x_{\text{imp}} \sim 20\,\text{nm}\). Simulations were used to evaluate \(x_{\text{imp}}\) as a function of field, and the results are plotted in figure 13.

5.2. Thermal meandering

The motion due to thermal fluctuations can be estimated using the Einstein equation,

\[
\langle x_{\text{thermal}}^2 \rangle = 2D\tau,\]

where \(x_{\text{thermal}}\) is the displacement in time \(\tau\) due to thermal motion and \(D\) is a diffusion constant. For one RF cycle at frequency \(f, \tau = f^{-1}\). \(D \approx \mu k_B T\), where \(k_B\) is Boltzmann’s constant, \(T\) is the temperature and \(\mu\) is the mobility of the vortex, given by Bardeen–Stephen as \(\mu = H_{c2}/(\phi_0 d)\), where \(\phi_0\) is the normal state resistivity, \(H_{c2}\) is the upper critical field and \(\phi_0\) is the flux quantum. Solving, the wandering due to thermal motion is given by:

\[
\sqrt{x_{\text{thermal}}^2} = \frac{2 k_B T \rho_n}{H_{c2} f \phi_0}.\]
For realistic parameters $T = 2 \text{ K}$, $\mu_0 = 100 \text{ n}\Omega\text{m}$; $\mu_0 H_{c2} = 30 \text{ T}$, $f = 1.3 \text{ GHz}$, $d = 30 \text{ nm}$, $x_{\text{thermal}} = 1.5 \text{ nm}$, as shown in figure 13. From these results, we can calculate the approximate expected rate of production of vortices that fail to annihilate. One expects that the distribution of final separations will be Gaussian with standard deviation $x_{\text{thermal}}$, suggesting that the number of vortices which do not annihilate will be given by the tail of the Gaussian. For example, at $H = 0.8 H_{c1}$, $x_{\text{imp}}$ is about 22 nm from figure 13, or about 15 standard deviations, making it extremely unlikely for vortices to escape due to thermal meandering alone.

5.3. Disorder meandering

To calculate the wandering due to surface disorder, illustrated in figure 14, we consider a single-cell $f = 1.3$ GHz niobium SRF cavity with an SIS structure using $d = 30 \text{ nm}$ thick Nb$_3$Sn layers. Assume that the topmost S layer has a normally-distributed random array of defects over its surface. For our geometry, we divide the $L \times L$ (where $L \sim 10 \text{ cm}$) surface area of the cavity into $N a \times a$ regions of order the pancake vortex size, where $N = L^2/\alpha^2$. We represent the effect of these defects as lowering the local value of $B_c$ in a given region. Therefore these defects will nucleate vortex penetration, and they will attract pancake vortices in the film. At the worst of the defects, the expected value for $H_c$ is $\alpha H_{c,\text{nominal}}$, where $\alpha$ is a constant between 0 and 1. At this defect, vortices penetrate at approximately $H = \alpha H_{c1}$. We represent the surface of the cavity with a distribution $\mathcal{H}$ of values for the reduction in the square of $H_c$. For simplicity of analysis, and since the defects are normally distributed, we will use the notation generally used in propagation of random errors.

$$\mathcal{H} = H_{c,\text{nominal}}^2 (1 - \alpha (0 \pm \sigma)).$$

Here is $\sigma$ the variance of the normally distributed $H_c^2$ reduction. We use absolute values because there should be no $H_c$ values higher than the nominal value.

We can find $\sigma$ using our restriction that the expected value for $H_c$ at the worst defect is $\alpha H_{c,\text{nominal}}$. To do this, we need to work with a normal distribution of the $H_c^2$ reduction in our $N$ regions $\mathcal{H}(0, \alpha) = \{1/(\sigma \sqrt{2\pi})\} \exp(\chi^2/2\sigma^2)$. First, we need to find $\phi$, the value of $x$ above which lies just one half of one of our $N$ regions (one half because the absolute value effectively doubles the number of samples in our integration).

$$\int_0^\infty \frac{1}{\sigma \sqrt{2\pi}} e^{-x^2/(2 \sigma^2)} \, dx = \frac{1}{2} \left(1 - \text{erf} \left( \frac{\phi}{\sqrt{2} \sigma} \right) \right) = \frac{1}{2N}. \tag{31}$$

Next, we set the expected value of the distribution in this region to be $1 - \alpha^2$. This sets the expected value of $H_c$ to be $\alpha H_{c,\text{nominal}}$ for the most extreme defect (we also have to normalize for there being only one defect in our sample size). This defines $\sigma$.

$$\int_0^\infty \frac{x}{\sigma \sqrt{2\pi}} e^{-x^2/(2 \sigma^2)} \, dx = \frac{\sigma}{\sqrt{2\pi}} e^{-x^2/(2 \sigma^2)} = \frac{1 - \alpha^2}{2N}. \tag{32}$$

To obtain an analytical expression, instead of solving equations (31) and (32), we can approximate. First, estimate that $\phi \approx \alpha$:

$$\int_0^\infty \frac{1}{\sigma \sqrt{2\pi}} e^{-x^2/(2 \sigma^2)} \, dx = \frac{1}{2N}. \tag{33}$$

We can also use a linear approximation for the Gaussian, so that the integral can be evaluated by calculating the area of the triangle shown in figure 15. The equation of the line evaluated at $y = 0$ determines the length $\Delta x$ in the figure:

$$0 = -\frac{\sigma^2}{\sqrt{2\pi}} e^{-x^2/(2 \sigma^2)} (\Delta x) + \frac{1}{\sigma \sqrt{2\pi}} e^{-x^2/(2 \sigma^2)}.$$  \tag{34}

This gives $\Delta x = \sigma^2/\alpha$. Evaluating the integral,

$$\int_0^\infty \frac{1}{\sigma \sqrt{2\pi}} e^{-x^2/(2 \sigma^2)} \, dx = \frac{\sigma}{2\alpha \sqrt{2\pi}} e^{-x^2/(2 \sigma^2)} = \frac{1}{2N}. \tag{35}$$

Rearranging,

$$\frac{\alpha^2}{2 \sigma^2} = \ln \frac{\sigma N}{\alpha \sqrt{2\pi}} \tag{36}$$

$N$ is very large, so $\ln \frac{\sigma N}{\alpha \sqrt{2\pi}} \approx \ln N$. Solving,

$$\sigma = \frac{\alpha}{\sqrt{2\ln N}}. \tag{37}$$

Now let us consider the behavior of the cavity at a field just above the expected vortex penetration field of the worst defect. Let $\hat{x}$ point into the film and $\hat{z}$ be aligned with the magnetic field. In this case, pairs of pancake vortices will form at the defect, move apart in $y$ due to the force exerted by the increasing magnetic field, and then move in the opposite direction in $y$ as the magnetic field direction reverses. In the mean time, they will have meandered some distance in $x$. If the meandering distance is very small compared to the impact parameter (approximately 50 nm), the pancake vortex pairs
are most likely to meet again and annihilate. However, if the meandering distance is significant compared to the impact parameter, vortex pairs are likely to get lost, accumulating over the film.

For this calculation, we will not do a full simulation. Rather, we will consider a path along $y$ of a vortex pancake moving from the defect at $y = 0$ to the extremum $y = y_{\text{max}}$ without any movement in $x$. We will, however, integrate the force in $x$ that the pancake would experience along its path, and calculate what the expected meandering distance would be from this.

The vortex will have two regions on either side of it in $x$ at a given time, one with $H_{c1}$, and one with $H_{c2}$. The centers of these two regions are separated by a distance $a$. The $x$-component of the force experienced by the vortex can be approximated from the gradient in condensation energy resulting from the difference in the local $H_c$. Magnetic energy density is given by $H^2/2\mu_0$. We convert this to energy by multiplying by the volume of a region, $a^2 d$.

\begin{equation}
F_x = \frac{H_{c1}^2 - H_{c2}^2}{2\mu_0 a^2 d}.
\end{equation}

The motion of the pancake is described by Bardeen–Stephen drag:

\begin{equation}
\eta \nu_x = F_x / d.
\end{equation}

Here $\nu_x$ is the $x$-component of the velocity of the pancake and $\eta$ is the Bardeen–Stephen drag coefficient, defined by $\eta = H_{c2} \phi_0 / \rho_n$, where $H_{c2}$ is the upper critical field of the film, $\phi_0$ is the flux quantum, and $\rho_n$ is the normal resistivity of the film.

The meandering distance $\Delta x$ within a region is given by $\Delta x = v \Delta t$, where $\Delta t$ is the time spent in that region. For simplicity, we approximate that the vortex moves with uniform speed in $y$, spending equal time in each of the regions that it travels through, such that $\Delta t = \frac{1}{2\nu_y}$.

Using equation (30) and equation (38), we can describe a distribution $F_x$ of forces that the pancake would experience. We observe that taking the difference between the absolute values from equation (30) produces a normal distribution:

\begin{equation}
F_x = \frac{a d H_{c,nominal}^2}{2\mu_0} (0 \pm \sigma).
\end{equation}

Between two regions that are adjacent in $x$, the pancake will experience a single value of the distribution of force values. After it travels a distance $a$, it will encounter a new pair of regions and therefore a new force value. We can use $\Delta x = v \Delta t$ and equation (39) to find $\sqrt{\langle \Delta x^2 \rangle}$, the rms wandering distance traveled by the vortex over a distance $a$:

\begin{equation}
\sqrt{\langle \Delta x^2 \rangle} = \sqrt{\langle F_x^2 \rangle} \frac{\Delta t}{\eta}.
\end{equation}

Multiply by the square root of the number of steps in a period $N_{\text{steps}} = \frac{2\nu_y}{a}$ to find the total rms wandering distance $\sqrt{\langle \Delta X^2 \rangle}$. Use equation (40).

\begin{equation}
\sqrt{\langle \Delta X^2 \rangle} = \frac{a d H_{c,nominal}}{2\mu_0} \frac{\Delta t}{\eta} \sqrt{\frac{2\nu_y}{a}}.
\end{equation}

Since $\Delta t = \frac{a}{\nu_y}$, from equation (39) and using $F_y = \frac{\phi_0 H}{\mu_0}$, we obtain:

\begin{equation}
\frac{\Delta t}{\eta} = \frac{a d}{F_y} = \frac{ad\mu_0}{\phi_0 \Delta H}.
\end{equation}

Here $\Delta H$ is the difference in magnetic field across the film. If we are looking just above the penetration field,
\[ \Delta H \approx \alpha H_d / \xi. \] Equation (43) can also be used to find \( \frac{\Delta \alpha \xi}{a} \).

\[
y_{\text{max}} = \frac{1}{2 \xi} = \frac{\phi_0 \alpha H_{\text{sh}}}{2 f_0 d \lambda \mu_0}.
\] (44)

Use the definition of \( \eta \).

\[
y_{\text{max}} = \frac{\rho_0 \alpha H_{\text{sh}}}{2 f H_{\text{c},2} a \lambda \mu_0}.
\] (45)

Now substitute.

\[
\sqrt{\langle \Delta X^2 \rangle} = \frac{1}{2^{\sqrt{\ln N}}} \frac{a^2 N}{\phi_0 H_{\text{sh}}} \left( \frac{\rho_0 \alpha H_{\text{sh}}}{f \xi H_{\text{c},2} a \lambda \mu_0} \right).
\] (46)

We then use \( H_{\text{c},\text{nominal}} = \frac{\phi_0}{2^{\sqrt{\ln N}}} \).

\[
\sqrt{\langle \Delta X^2 \rangle} = \frac{\xi}{4 \sqrt{2 \pi \ln N}} \frac{H_{\text{c},\text{nominal}}}{H_{\text{sh}}} \left( \frac{\rho_0 \alpha H_{\text{sh}}}{f \xi H_{\text{c},2} a \lambda \mu_0} \right) .
\] (47)

We can set our region size \( a = \xi \approx 4 \text{ nm} \). \( H_{\text{sh}} / H_{\text{c},\text{nominal}} \) for Nb\(_3\)Sn is approximately 0.75. \( N \) is approximately \( 10^{12} \), so \( 1 / (4 \sqrt{2 \pi \ln N}) \) is approximately \( 10^{-2} \). We also use \( H_{\text{c},2} = 30 \text{ T} \), \( \rho_0 = 10 \text{ n}\Omega \text{m} \), and \( H_{\text{c},\text{nominal}} = 530 \text{ mT} \). For \( \alpha = 0.8 \), these factors give \( \sqrt{\langle \Delta X^2 \rangle} \approx 2.1 \text{ nm} \). Setting this as a standard deviation for vortex meandering and using \( x_{\text{imp}} \approx 22 \text{ nm} \) from above gives a result of 11 standard deviations, again making it extremely unlikely for vortices to escape due to disorder meandering.

Varying \( a \) and \( \alpha \) over a reasonable range also gives a small value for the meandering distance, as seen in figure 16.

In addition to the impact parameter and thermal meandering distance, figure 13 plots the meandering distance due to disorder using the analytical formulation above. Also plotted in the figure is a simulation of vortex pair creation and annihilation in the presence of disorder. In addition to forces from the applied magnetic field and from the randomly distributed array of circular defects with radius \( a \)\(^{17}\), the simulation considers the forces of one vortex on the other, and finds the maximum lateral separation between the pair during a cycle.

5.4. Interacting vortices, interacting defects

We have presented analyses of thermal meandering and meandering of a vortex–antivortex pair due to disorder, and so far, nothing has caused a large buildup of vortices in the film. We have also performed a preliminary analysis of the nucleation of many vortices at a defect over the course of an RF cycle, all of which interact with each other. While an attractive force exists between vortices and antivortices, vortices exert a repulsive force on other vortices, and antivortices exert a repulsive force on other antivortices. These repulsive forces between similar vortex types appear to result in substantially larger lateral movement, which could lead to higher rates of non-annihilation. An example video is included as supplemental material [86]; three frames of which are shown in figure 17. Here the film thickness is 30nm, parameters are appropriate for Nb\(_3\)Sn material, and the disorder is modeled as 60 pinning sites randomly spread over 0.4

\(^{17}\) In the simulation, the defects pin vortices with force that increases linearly from the defect edge where it is zero, to the center of the defect, where it is a maximum. The maximum is set such that moving the vortex from the center of the defect to the edge would require work equal to the condensation energy of the volume of the vortex core.
square microns that exert a force of $F_{\text{max}} = k_i \rho$, where $F_{\text{max}} = B_i^2 \rho \xi^2 \frac{d}{r_{\text{pin}}} / 2$, $k_i$ is a value between 0 and 1 that is randomly chosen for each pinning site, and $\rho = r_{\text{sep}} / r_{\text{pin}}$ for $r_{\text{sep}} < r_{\text{pin}}$ and 0 for $r_{\text{sep}} > r_{\text{pin}}$, where $r_{\text{sep}}$ is the distance between the vortex and the center of the pinning site and $r_{\text{pin}}$ is the pinning site radius, (here chosen to be 2 nm). Note that the maximum horizontal meandering due to disorder and interactions is roughly eight nanometers, only a factor of three less than the impact parameter $x_{\text{sep}}$, suggesting that interactions may be much more dangerous for multiple defects over many cycles. However, further work on interactions is clearly needed; it is possible that realistic parameters for the disorder strength could decrease the meandering, and it is possible that the distribution of maximum meandering distances near a defect over multiple cycles could be sub-Gaussian. On the other hand, it is likely that two defects that both nucleate pancake vortices and are close enough together that the vortices interact could be dangerous even in the absence of disorder.

6. Conclusions

This article attempts to make a case for thoughtful, broad efforts to identify and estimate fundamental physical limits of materials in parallel with experimental efforts. Our estimates for the superheating field of pure materials [14, 15] had a significant impact in the SRF community, we understand, not because it promised that Nb$_3$Sn could be run at twice the field of Nb. Rather, we pointed out that the ‘vortex line nucleation’ model was incorrect (see footnote 9). This model, created by and initially trusted in the SRF community, had provided discouraging estimates for high $\kappa$ materials, misleading the field for some years. We also note that our controlled, theoretically grounded calculations allow one to explore questions like materials anisotropy that cannot be gleaned reliably from phenomenological models (or, indeed, from our own qualitative arguments, section 4.2).

We understand that many in the SRF community are skeptical of the use of bulk (or thick films) of new materials with higher $\kappa$, even though the theoretical estimates suggest significantly improved $H_{\text{th}}$ as well as lower cooling requirements. We too were concerned until recently that the smaller coherence lengths might make the metastable state more susceptible to vortex penetration. But we believe that our calculation of the effects of disorder within a particular model clearly indicates that the reliability of the new materials increases so rapidly away from $H_{\text{th}}$ that the effects of lower coherence length should not be a major concern. One must always make choices of where to focus resources (laminates versus bulk materials, coated copper versus niobium [87]; an interesting review has been recently published in this special issue [74])—but informed choices may involve consultation with experts outside the SRF field.

We are also guardedly pessimistic about the utility of thin laminates in increasing performance. (a) We are concerned that experimentalists continue to be inspired by the very high parallel fields sustainable by isolated thin films [74, section 9] (see footnote 15), even though these high fields are irrelevant in GHz applications [76] and likely also practically inaccessible in any AC application. (b) Dangerous vortices in thin films will not typically reside parallel and inside the films, but penetrate in and out of the film via pancake vortices, whose motion dissipates heat. The maximum fields reachable without flux penetration for ideal thin films are rather similar to those of the bulk material [45]. The flux penetration needed to reach higher fields produces heating per cycle comparable to that for copper cavities [76]. (c) We explore extensively the suggestion [73] that the insulating layers in laminates may act to trap flux from defects, keeping flux from entering the bulk. Here the key issue is whether the nucleated pancake vortices escape from the vicinity of their parent flaw, or annihilate with their partners at the end of each cycle. Modeling these as sources for pancake vortices, we agree that neither thermal diffusion nor disorder are dangerous, but that interactions between vortices, and between vortices generated at separate defects, might allow for escape and unacceptable heating—warranting experimental caution and further theoretical study.

Finally, a word to our colleagues outside the accelerator community. This paper is a collaboration of SRF experts from the accelerator community (Posen, Liepe) and condensed matter theorists, and we draw heavily on conversations with both theorists and experimentalists inside the community (Hasan Padamsee, Alex Gurevich, Nicholas Valles, Takayuki Kubo, Kenji Saito). These domain-specific experts have enormous experience in the challenges and issues relevant for the field; we were told that superheating fields, higher $\kappa$ materials, anisotropy, disorder, and laminates were the key questions, and have been guided into studying these in the correct limits and focusing on the right issues. We can testify that this teamwork has led to both excellent condensed-matter physics and efficient, targeted research to improve SRF performance.

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