How Solid is Supersolid?

Anatoly B. Kuklov
Department of Engineering Science and Physics, CSI, CUNY, Staten Island, NY 10314, USA

Nikolay V. Prokof’ev and Boris V. Svistunov
Department of Physics, University of Massachusetts, Amherst, MA 01003, USA and Russian Research Center “Kurchatov Institute,” 123182 Moscow, Russia

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Macroscopic quantum properties of helium-4, one of the simplest and oldest elements in the universe, continue to puzzle and amaze scientists. Supertransport in solid helium-4 is the most elusive and controversial conundrum of all.

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Among known states of natural matter, supersolids are often regarded as counterintuitive in the extreme. For more than 50 years, a number of authors with different perspectives such as Gross [1], Andreev and Lifshitz [2], Thouless [3], and Chester [4], have contemplated this exotic state. Thought to be the coexistence of superfluid and crystalline orders in the same single-component material, supersolids still attract broad interest. The original predictions were made for perfect quantum crystals in free space, with the most promising candidate being solid helium-4 ($^4$He), which is known to have the largest amplitude of zero-point motion of atoms in the ground state. Such motion was presumed to create a stable (repulsive) gas of zero-point vacancies undergoing Bose-Einstein condensation at low temperature. Thus the perfect crystal would simultaneously be a solid and a superfluid. This simple and attractive idea sparked a wide experimental search for supersolidity in solid $^4$He, which, however, yielded no positive results. Recent interest in the supersolid state of matter was rekindled after observations of a signal consistent with supersolidity in the torsional oscillator experiment performed by Kim and Chan at Penn State [5] in 2004. In what follows, we will survey both the recent experimental findings and the theoretical landscape in supersolid research.

In terms of theory and calculations, the problem of supersolidity in free space for solids with one or two atoms in the unit cell is not amenable to any existing analytical scheme. This is the result of strong competition between the two orders—superfluid and crystalline—at the interatomic distance. In fact, the crystalline lattice is free to adjust its period continuously in order to eliminate vacancies and/or interstitials introduced out of equilibrium. As of today, there is no clear understanding of what is required to obtain a stable supersolid state in free space with few atoms in the unit cell, and this is one interesting research direction to explore.

Widespread efforts are also aimed at finding simplified models of supersolids. Lattice supersolids are quite well studied nowadays [6], in such systems the lattice is imposed externally, and accordingly, the solid period is forced to take a discrete set of values, i.e., it cannot change continuously. Crystalline order in such lattice models breaks the discrete translation invariance of the external lattice potential and is typically “pinned” to the lattice, as shown in Fig. 1. As a result, excitations of such order cost finite energy—as opposed to a solid in free space. The superfluid component can be introduced by doping the solid, i.e., by adding (removing) atoms to produce particle (hole) excitations that, if stable against phase separation—which is always an option—Bose-condense. In many respects, this picture is similar to that of the so-called doped Mott insulator. In such an insulator, the number of particles is exactly equal to the number of the lattice sites. Should the onsite repulsion between particles significantly exceed the kinetic energy loss due to localization, particles become localized—one at each site. If, however, the balance between particles and sites is violated, the insulator becomes a superfluid. Most importantly, pinned solid structures lack the ability to adjust their period continuously to make it commensurate with the particle number and eliminate the superfluid component (shown by open circles in Fig. 1). It seems that most known lattice supersolids are of this kind, with a notable exception presented recently in Ref. [7].

At the other end of the problem we find Bose-Einstein condensates with the spatial modulation of density induced by interparticle interactions, as first envisioned by Gross [1] (see also Ref. [8]). These supersolids are best described within the classical field approach, relying on the mean-field description of bosons in terms of the so-called Gross-Pitaevskii equation, where the quantum bosonic operator is replaced by a classical field. They emerge in the special limit of weak interatomic potentials and a large (infinite in the classical-field limit) number of particles in the unit cell. A nontrivial situation occurs when the spatial period becomes comparable with the interparticle distance. In this regime, the discreteness of particle number becomes important and the Gross-Pitaevskii approach fails when this number approaches unity.

One may also think of tailoring a two-body potential...
FIG. 1: An example of the lattice supersolid: the checkerboard order formed by particles (filled circles) in the underlying square lattice coexists with the superfluidity of mobile vacancies (open circles). (APS/A. B. Kuklov et al.)

to extend the region of stability for the supersolid phase in free space, starting from the classical field perspective. This idea was realized by Cinti et al. and Sacconi et al.[9], when a microscopic model featuring the supersolid phase (with the number of particles per self-organized unit cell being around 10) was constructed. The potential consists of a generic repulsive part, saturating to a constant below some radius $R_0$. The microscopic structure of the supersolid state is that of liquid droplets (each of a radius $\sim R_0$) arranged in a regular solidlike structure. Since particles in each droplet are in a liquid state, the overall phase coherence is achieved by means of Josephson coupling between the droplets. Since the interatomic distance is much smaller than a typical wavelength of the density modulations, the discreetness of the particle number in the unit cell becomes irrelevant and the resulting supersolid can be described within the classical field approach, that is, following from the solutions of the mean-field equations. These key features are depicted in Fig. 2, which shows the shapes of the interaction potentials used in Refs. [8, 9], as well as the resulting density modulation—the crystalline-type order. It would be interesting to establish for some reference cases where the boundary between the classical and quantum worlds stands in terms of the number of particles per unit cell. At the moment it seems very unlikely, though not improbable, that a supersolid with one atom per unit cell exists for a two-body interaction potential of a standard shape [10]. Similar classical field considerations apply to lattice models in the limit of large occupation numbers and weak interactions (on-site and nearest-neighbor), ensuring large supersolid regions in the parameter space.

FIG. 2: Two-body interaction potentials (the upper panel)—$V_{\text{soft-core}}(r)$ [9] and $V_{\text{roton-type}}(r)$ [8]—leading to a supersolid characterized by spatial density modulations (shown in the lower panel) significantly larger than the interparticle distance. Vertical axes—arbitrary units. Horizontal axes—distances in units of the interparticle separation. (APS/A. B. Kuklov et al.)

Remarkably, the supersolid behavior in such lattice models survives even at filling factors close to $1/2$[7]. Although not yet observed experimentally or numerically, the supersolid state is guaranteed to exist in two-dimensional dipolar systems. This is an immediate consequence of the theorem that first-order transitions that involve a density change, such as liquid-solid ones, are forbidden due to log-divergent negative terms in the surface tension (see, e.g., Ref. [12]). It is thus guaranteed that on approach to the solid phase from the superfluid side of the phase diagram (by changing any control parameter) one can construct a state containing large solid domains that is lower in energy than the homogeneous liquid. At $T = 0$, such domains will form a secondary lattice superstructure leading to the global supersolid phase (in fact, a whole set of supersolid phases [12]). The microscopic picture of this phenomenon, however, was never quantified by measuring the surface tension parameters.

The experimental situation in solid $^4$He

Apart from general fundamental questions, the main puzzle is to understand what happens in $^4$He, where both orders are in strong competition with each other. Initial attempts to detect supersolid properties [13] have yielded essentially nothing, including the new results in solids subject to pressure gradients [11]. However, follow-

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ing Leggett’s torsion oscillator (TO) proposal [15] and hints of anomalous behavior found by Goodkind in ultrasound measurements of hexagonal-close-packed (hcp) \(^4\)He at temperatures \(T \approx 0.2\ \text{K} [16]\), in 2004 Kim and Chan discovered a change in the TO period. They interpreted it as the signature of the nonclassical rotational inertia (NCRIF) of \(^4\)He atoms in the solid state \([5]\). Since then, small TO period reduction by a few nanoseconds (for a typical resonant period of \(\sim 1\) millisecond) below \(T \approx 0.2\ \text{K}\) has been reproduced by many groups all around the world. Specific heat measurements also show an anomaly (a wide and weak peak) in the same temperature range \([17]\). It is also worth mentioning quite unique setups detecting the TO effect under overall dc rotation \([18]\). Some observed features, however, (such as low-\(T\) dissipation) are still controversial and require further experimental investigation.

There are numerous difficulties with the standard suprasolid interpretation of the period decrease. The most striking ones are as follows:

(i) The NCRIF fraction (NCRIF) is very sensitive to the TO cell design: the NCRIF of \(\sim 1\% [5]\) originally detected in a composite cell was found either to drop by \(\sim 100\) times in a much more rigid cell \([19]\) or increase by a factor of \(20\) in a narrow annulus space \([20]\). Polycrystalline samples grown by the blocked capillary technique may have both larger and smaller NCRIF than samples grown at constant temperature from the superfluid, not to mention large annealing effects \([20]\).

(ii) The potential flow tests (verifying that NCRIF is sensitive to changes in the topology of the flow pattern) remain controversial \([21]\).

(iii) There is no clear evidence of a phase transition yet. Typically, it is a crossover taking place within the temperature interval \(0.1\ \text{K} < T < 0.25\ \text{K}\), which is barely changing—despite enormous variations in the NCRIF by several orders of magnitude (!)—from one TO setup to the next. This is not expected even in strongly disordered 3D superfluids.

(iv) An enigmatic feature, which is impossible to reconcile with the uniform condensation of vacancies in a 3D sample, is a dissipation peak centered at the temperature of the strongest variation of NCRIF \([5]\). This peak shows up as a dip in the amplitude of the resonant oscillations vs temperature.

(v) The NCRIF suppression, occurring when the rim velocity \(v\) oscillates at an amplitude as small as \(v \sim 10–100\ \mu\text{s} / \text{s}\), cannot be accounted for by any known mechanism of lowering the superfluid critical velocity from a typical value of a few m/s. More importantly, the effect of velocity does not seem to be of a sharp threshold character, with a pronounced critical value \(v_c\). Rather, it is found to be equivalent, in a certain sense, to temperature \([22]\): the NCRIF depends on a single variable \(\xi \sim (T/T_0)^\lambda + (v/v_0)^\lambda\), with \(T_0, v_0, \xi > 0, \lambda > 0\), \(\lambda/\xi \approx 0.43\) being some parameters. Such collapse of the data into a single master curve, in particular, implies that \(v_c\) must be an increasing function of temperature: \(v_c \sim T^\xi/\lambda\), with \(\xi/\lambda \approx 2.3\). However, this feature has been recently questioned by Kojima whose data suggests that \(v_c\) first significantly decreases with \(T\) and then saturates to a constant value \([23]\).

(vi) At low temperature \(T_h < 50–70\ \text{mK}\), the NCRIF data demonstrate hysteretic behavior as a function of velocity \([24]\). To some extent, a summary of this feature and of the previous one can be presented in the form of a “phase diagram” in the \((v, T)\) plane \([25]\).

(vii) Extremely small concentrations of \(^3\)He impurities (as small as 10 ppb–1 ppm) strongly affect the onset temperature and the NCRIF amplitude \([26]\).

The phenomena described above (and also the results of first-principles simulations discussed below) unambiguously indicate that NCRIF is induced by structural disorder. Nevertheless, a consistent microscopic picture behind the experimental findings is still missing. Being the hallmark signature of the superfluid behavior, sensitivity to the sample topology has to be settled beyond reasonable doubt in future experiments. The current research trend is to design new experiments to attack the problem from different angles; on the theoretical side, the challenge is to understand the mechanisms leading to the observed striking phenomena. Below we review some of the new and existing directions.

A strong indication of structural disorder comes from NMR experiments \([27]\) where a large fraction of \(^3\)He impurities (about 100 ppm) is trapped by some lattice defects. If these defects are dislocations, then their density is at least 4 orders of magnitude larger than the traditionally accepted values in the bulk. Most recent NMR measurements \([28]\) at low \(^3\)He concentrations have found a peak in \(T_1\) relaxation time (a measure of how fast the magnetization along the applied magnetic field relaxes to its equilibrium value) at temperatures where the TO anomaly, specific heat anomaly \([17]\), and the maximum of magnetization along the applied magnetic field relaxes to its equilibrium value) at temperatures where the TO anomaly, specific heat anomaly \([17]\), and the maximum of dissipation are observed \([5]\). This peak \([28]\) might be an indication of some phase transformation occurring within the defect structure.

An unexpected discovery by Day and Beamish \([29]\) was that the shear modulus \(\mu(T)\) of hcp \(^4\)He hardens below \(T_0 \approx 0.15\ \text{K}\) at frequencies 2–2000 Hz—exactly in the range of the TO anomaly. Such a hardening in solid \(^4\)He was observed much earlier by Tsymbalenko \([30]\) at significantly larger temperatures and frequencies. At the moment it is not clear if these two anomalies are due to one and the same mechanism. Originally, the low-temperature hardening effect was observed by Friedel \([31]\) in pure aluminum more than 50 years ago. Amazingly, many properties of \(\mu(T)\) and NCRIF regarding temperature and rim velocity dependencies, the responses to \(^3\)He impurities, or the hysteretic behavior \([24, 29]\) are similar, if not identical, to each other. The hardening of \(^4\)He can be related to pinning of dislocations by \(^3\)He impurities \([29]\) as well as to self-trapping \([32]\) in the periodic potential provided by the surrounding crystalline lattice—the so-called Peterle potential. Within the linear response regime, the shear modulus increase leads to the
TO period decrease at frequencies well below acoustic resonances; i.e., there is a possibility for a purely “classical” interpretation of the TO anomaly. This explanation, however, does not pass the quantitative test: numerical evaluation of the TO period shift due to a 20% change in $\mu$ shows that it is barely a few percent of the observed effect. The other difficulty is that similar period shifts, including hysteresis, small critical velocities, and the effect of small concentrations of $^3$He are observed in porous materials (Vycor, porous gold, and Gel-sil) where dislocations, whatever they are in such highly disordered medium, should be strongly pinned by pore walls. Finally, critical stresses for suppression of the shear modulus effect are almost 2 orders of magnitude larger than stresses generated by ac velocities $\sim 10 \, \mu m/s$ influencing the NCRI effect.

The “ordinary glass” approach involves proposing a phenomenological model for the generalized response function to describe period shifts using several fitting parameters and functions. The goal of this phenomenology is to demonstrate that the experimentally observed response is consistent with zero superfluid fraction and thus can be of nonsuperfluid origin. This approach, however, does not explain the microscopic origins of the “ordinary glassiness,” which is known not to exist (at least, on the classical level) in solids made of simple atomic elements, nor does it explain the physics behind model parameters and their temperature dependencies. In this sense the ordinary glass framework is as enigmatic as the supersolid interpretation itself.

Recent work by Reppy posed serious experimental objections to the idea of a superfluid origin of the period shift. In this experiment, the change of the period of oscillation—as a function of deliberately introduced disorder—was observed at high ($\sim 1 \, K$), not low, temperatures, while in the $T \to 0$ limit the oscillation periods of samples with different amounts of disorder were approaching one and the same value. Similar behavior was observed upon annealing—period changes occurred at high $T$. Clearly, experiments of the kind carried out by Reppy (or at least the annealing part) should be repeated by other groups to settle the issue. The most recent results on the triple-frequency TO have clearly demonstrated that the NCRI can be extremely sensitive to the mechanical properties depending on the TO design. The possibility of the nonsuperfluid origin of the NCRI was also reported by Eyal et al., where a strong NCRI effect has been seen at temperatures as high as $1.3 – 1.9 \, K$—well above the 0.2 K anomaly.

Given the rather unknown character of structural disorder in solid $^4$He, it is desirable to study behavior of almost ideal crystals. Working in this direction, the group at École Normale Supérieure in Paris is now growing essentially monocrystalline $^4$He samples, free from $^3$He impurities, and is performing ultrasound and TO experiments on them. One of the striking effects found recently is the strong low-$T$ shear modulus softening effect—just the opposite to the observation of Day and Beamish. Such softening is metastable—it disappears upon the thermal cycling. The nature of this effect requires further study.

### Observation of direct superflow through $^4$He

A unique experimental setup for studying the dc flow response of solid helium was created by Ray and Hallock. In these experiments, atoms of $^4$He can be injected into solid $^4$He from the superfluid reservoirs through two Vycor rods. A schematic of the setup is shown in Fig. 3, the so-called syringe regime, when both electrodes are utilized for injecting liquid helium into the solid. In flow experiments one looks at the system response when pressure/temperature is changed in one reservoir by monitoring pressure/temperature in the second reservoir. The syringe regime is used for studying the so-called isochoric compressibility, that is, the variation of the density of atoms— injected one-by-one into a constant-volume solid phase—in response to varying the outside chemical potential. Such compressibility has been found to be comparable to that of liquid $^4$He, despite the fact that in an ideal solid it is essentially zero. As suggested by Söyler et al., crystal growth is possible by the mechanism of dislocation superclimb—edge dislocations add atoms to atomic layers provided there is superfluid atomic transport along their cores (see Fig. 3). It is important that the giant isochoric compressibility regime has always been observed together with the superflow through the solid. Furthermore, while the solid may contain frozen-in pressure gradients of about 0.1 bar, the accumulation of matter under the injection occurs in a uniform manner—the pressure rises in unison at the opposite ends of the sample. This feature is clearly inconsistent with any type of classical plastic behavior, which results in time-dependent pressure gradients. At the same time, the picture of a percolating network of dislocations with superfluid cores delivering matter independently of the pressure gradients is consistent with such an observation.

Another important piece of evidence for superfluidity in the experiment is the observed regime of supercritical flow—linear dependence on time of the pressure difference between the two superfluid reservoirs. The value of the onset temperature $T \approx 0.5 \, K$ is about 4 times smaller than the $\lambda$ temperature of liquid $^4$He. This excludes the interpretation that the flow proceeds along stable mesoscopic liquid channels, which are known to exist in polycrystalline samples at the melting curve. The strongest evidence against the liquid-channel picture is a striking and dramatic drop of the critical dc flux taking place at temperatures around 70 mK, with a surprising subsequent revival at temperatures around 60 mK. This phenomenon appears to be the key to understanding properties of superfluid structural defects in solid He.
FIG. 3: Schematic diagram for the Ray and Hallock setup depicting the superclimb effect. The main rectangle features solid $^4$He, containing an edge dislocation with the superfluid core shown as the thick blue line. Dashed lines outline the Vycor “electrodes” protruding from the liquid $^4$He reservoirs (rectangles atop of the Vycor) into the solid. The “electrodes” feed superfluid $^4$He (shown in blue) from the reservoirs to the dislocation core. As the level of superfluid in the reservoirs decreases (shown by the grey arrows), the core climbs upward (in the direction of the red arrow) which results in building up the extra plane of atoms shown by red hatching. (APS/A. B. Kuklov et al.)

and definitely deserves more attention.

Given the extremely controversial character of the TO data in terms of their superfluid interpretation, the natural question to ask is whether phenomena revealed by Ray and Hallock have anything to do with the TO behavior. Formally, the results by Ray and Hallock do imply that there should be some mass decoupling corresponding to the superfluid fraction. However, the estimates of the superfluid fraction they observed show that it is too small to be resolved in a TO experiment. An intriguing connection might be through yet-to-be-understood quantum physics of structural defects.

**Theoretical understanding of solid $^4$He**

Theorists have also been hard at work shedding light on what is and what is not possible in bosonic continuous-space solids with the typical interparticle interaction $U(r)$, and in particular, in the hcp crystal of $^4$He. Let us consider an interaction potential that does not favor vacancies or interstitials in the classical, not quantum, crystal at zero $T$ (in fact, we are not aware of any counterexample) and ask whether quantum mechanics can change this situation and induce zero-point vacancies or interstitials (to become superfluid according to the Andreev-Lifshitz scenario). Remember that Feynman’s path integral representation of the equilibrium statistics of bosons in $d$ dimensions establishes an exact mapping onto statistics of classical $(d + 1)$-dimensional “polymers” (particle world lines). With the celebrated Pollock and Ceperley formula this yields an explicit relation between the variation of polymer windings and superfluid response. Thus supersolidity can be pictured as a disordered state of world lines of vacancies (interstitials) among relatively straight and ordered world lines of the rest of the bosons.

Within the path integral representation one can perform first-principles simulations of systems consisting of

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thousands of $^4$He atoms. This representation is also a powerful tool for general qualitative analysis. Using the language of path integrals, Prokof’ev and Svistunov have proven a theorem (see Ref. [26] in Ref. [46]) that no supersolid is possible without such zero-point defects. The crucial control parameter $\xi_{\text{ins}} = mU/\hbar^2$, where $m$ denotes the particle mass, and $U$ and $R_0$ are the characteristic values of the interaction potential and interparticle distance, respectively. This parameter controls the ratio between the potential and zero-point kinetic energy terms for vacancy/interstitial world lines. Large $\xi_{\text{ins}}$ ensures that no zero-point vacancies (or interstitials) can be induced. In other words, for large $\xi_{\text{ins}}$ the crystal adjusts its volume so that no zero-point defects—either vacancies or interstitials—are present in the ground state. Furthermore, energy gaps for their creation in pairs are large (cf. the discussion presented by Andreev and Lifschitz in Ref. [2]), and thus the ground state is insulating according to the above mentioned theorem.

Reliable first principles Monte Carlo simulations allow us to explore basic properties of ideal hcp crystals, as well as point and extended defects embedded in it. These simulations have demonstrated that a perfect hcp crystal of $^4$He is an insulator with the gap of about 13 K for vacancies and 23 K for interstitials (see Refs. [32,33] in Ref. [46]). Furthermore, it has been numerically observed that injection of vacancies into a perfect sample leads to their clustering (see Ref. [38] in Ref. [46]) into Frank’s dislocation loops (see Ref. [31]) in the basal plane (or phase separation at the melting curve). These results invalidate the Andreev-Lifshitz-Thouless-Chester (ALTC) proposal.

Having argued against the ALTC supersolid, we stress that the hcp solid $^4$He is likely to support supersolidity induced by a network of superfluid structural topological defects. $Ab$ initio Monte Carlo simulations have revealed that some grain boundaries [37] and dislocations [44,48] support superflow. More than two decades ago, Shevchenko predicted very unusual dynamical properties of a network of superfluid dislocations [49]—large temperature range where at extremely low frequencies the superfluidity of their cores as it requires mass transport in the crystal. Most recently, it was noted [53] that the dip in the flow rate [54] may be due to the dislocation roughening induced by the frozen internal stresses. Superclimb is one of the qualitatively new effects in the emerging subfield of metallurgical behavior induced by quantum phenomena—quantum metallurgy,” as called by A. Dorsey. It provides strong evidence that supersolidity of structural defects and metallurgical behavior of solid $^4$He are closely interconnected and the setup of Ray and Hallock [53] is highly promising for studying this new physics.

Solid $^4$He may also exist in the metastable superglass state observed in Monte Carlo simulations of quenched liquid $^4$He (see Ref. [32] in Ref. [46]). This state is characterized by the absence of any long-range structural order and superfluid fractions as large as ten percent. It is, however, unlikely that this phase can form under the experimental conditions of relatively slow cooling in TO. A recent experiment [52] challenges this view and suggests that bosonic $^4$He, as opposed to fermionic $^3$He, can form stable glass or dense dislocation network at low $T$. It has been observed that inducing structural disorder by external stress in solid $^4$He leads to the $\sim T^2$ contribution to pressure. No such effect has been seen in samples of solid $^3$He. This contribution is consistent with linear specific heat due to two-level systems in glasses and/or dislocations.

**New challenges**

Clearly, more measurements are needed. In our opinion, in order of priority, it is crucial to measure the isochoric compressibility $\chi(T,\omega)$ and the critical superflow rate $R(T,\omega)$ as functions of temperature $T$, frequency $\omega$, and...
and $^3$He concentration. The superclimb effect is expected to be suppressed below a temperature $T_r$, determining dislocation roughening. Such suppression must lead to reduced $\chi(T, \omega)$ and enhanced $R(T, \omega)$. We note that $R(T, \omega)$ and $\chi(T, \omega)$ are sensitive to the nature of the Shevchenko state, that is, to the phase-slip rate. Such dependencies will provide valuable information for constructing theoretical models of the network where purely superfluid dynamics is coupled to climbing and gliding motion of structural defects. One possible structure of such dynamical network is shown in Fig. 5. Another important aspect is understanding the nature of the dip in the flow rate at low temperature and, in particular, its shape and position as a function of $^3$He concentration (see the proposal in Ref. [53]). The “smoking gun” evidence for supersolid behavior would be the observation of the persistent currents.

The Ray and Hallock setup can shed light on the fundamental question: Why was there no flow observed upon applying mechanical pressure to the solid in a number of experiments [13, 14]? In this regard we note that the syringe effect immediately implies the possibility of reversing it, i.e., creating a flow of matter from the solid to a reservoir (via a super link) upon applying pressure to the solid. To what extent the very large aspect ratio (length/width $\sim 100–400$) of the capillaries and, possibly, the crystal growth conditions in the setups [13, 14] are relevant to the answer remains to be seen. Interestingly, Ray and Hallock have observed that the syringe effect does not take place if the crystal is grown by injecting atoms into the superfluid at sufficiently low temperature [43].

The “conductance through solid” measurements can be helpful in resolving yet another puzzle, namely, the close similarity between all TO features observed in crystalline $^4$He and solid $^4$He confined in Vycor [5] and other nanopore materials [34]. The key question is whether or not there are differences in $R(T, \omega)$ and $\chi(T, \omega)$ in both media. At this point it is instructive to recall the history of discovering linear topological defects in solids—the dislocations and disclinations. This discovery was made by Vito Volterra long before the crystalline structure of solids had been seen [55]. In fact, the only requirement for having a dislocation characterized by typical long-range stresses around the core is that a medium can sustain shear stresses up to some threshold. Thus, while crystallinity introduces exact quantization of the Burgers vector and related topological protection of the defect, such exactness is not required for characterizing dislocations as linear concentrators of stresses. Dislocations introduced at this level of description could explain the similarity between TO responses of crystalline $^4$He and strongly disordered solid $^4$He in Vycor.

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\text{FIG. 5: The structure of the percolating Shevchenko network of superfluid dislocations can, in principle, be very complicated (see top panel—a “plumber’s nightmare”). The schematic in the bottom panel shows one of the simplest possibilities: Vertical lines represent superfluid screw dislocations linked to the superfluid and superclimbing edge dislocations forming prismatic loops (shown by elliptical shapes). The inside of such loops are the (insulating) basal plane fcc faults. The links between the dislocations are shown as blue solid dots. The percolation of the superflow occurs in the vertical (along the hcp axis) as well as in the horizontal (along the basal plane) directions. One such path is shown by the dotted green line. ((Top) iStockphoto.com/ilbusca; (Bottom) APS/A. B. Kuklov et al.)}
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References

About the Authors

Anatoly B. Kuklov

Anatoly Kuklov received his Ph.D. from Odessa State University, Ukraine, in 1986. He worked in the Research Institute of Physics, Odessa, Ukraine, 1980–1992 and joined the City University of New York (CUNY) in 1993. Currently he is a professor of physics at the College of Staten Island, CUNY, and at the Graduate Center of CUNY. His research focuses on strongly correlated systems, which include supersolids, superfluids, superconductors, and exotic phases of ultracold atoms.

Nikolay V. Prokof’ev

Nikolay Prokof’ev received his Ph.D. in 1987, from Kurchatov Institute, Moscow, where he worked from 1982 to 1999, with a period between 1992 and 1994 spent at the University of British Columbia, Vancouver, as an International Fellow. In 1999 he joined the Physics Department of the University of Massachusetts, Amherst. His research deals with strongly correlated states in electronic and bosonic systems, critical phenomena, and quantum Monte Carlo methods. He is a Fellow and Outstanding Referee of the American Physical Society.

Boris V. Svistunov

Boris Svistunov received his Ph.D. in 1990, from Kurchatov Institute (Moscow) where he worked from 1986 to 2003 (and which he is still affiliated with). In 2003 he joined the Physics Department of the University of Massachusetts, Amherst. His research deals with ultracold gases, superfluidity and supersolidity, strongly correlated systems, and the theory and practice of quantum Monte Carlo methods. He is a Fellow and Outstanding Referee of the American Physical Society.