1	Multiple Coulomb phase in the fluoride pyrochlore CsNiCrF ₆
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15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38	The Coulomb phase is an idealized state of matter, whose properties are determined by factors beyond conventional considerations of symmetry, including global topology, conservation laws and emergent order. Theoretically, Coulomb phases occur in ice-type systems like ice and spin ice; in dimer models; and certain spin liquids. However, apart from ice-type systems, more general experimental examples are very scarce. Here we study the partly-disordered material CsNiCrF ₆ and show that this material is a multiple Coulomb phase with signature correlations in three degrees of freedom: charge configurations, atom displacements, and spin configurations. We use neutron and x-ray scattering to separate these correlations, and to determine the magnetic excitation spectrum. Our results show how the structural and magnetic properties of apparently disordered materials may inherit, and be dictated by, a hidden symmetry – the local gauge symmetry of an underlying Coulomb phase.
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44 The correlations and dynamics of conventional phases of matter can be summarized in terms 45 of broken symmetries [1] and the transverse (Goldstone) and longitudinal (Higgs) fluctuations of their associated order parameters [2]. By contrast, in a Coulomb phase [3], there is no long-46 47 range order and the cooperative behavior of the local degrees of freedom is best described by 48 a field, whose emergent symmetry is that of the electromagnetic field. Any ground state of the 49 local degrees of freedom can be represented by a non-divergent field (i.e. div**B**=0, see Fig. 1A), 50 which means that although the Coulomb phase has no broken global symmetry, it has local 51 gauge symmetry and the closed loop topology of the associated field lines [3,4,5]. Dynamics 52 involve coherent fluctuations of the field, or topological defects in the field, which can 53 respectively be identified as generalized photons and charges of the relevant field theories 54 [6,7,8]. 55

- 56 Although the charge ice [9,10] (Fig. 1B) and the pyrochlore Heisenberg antiferromagnet [11] 57 (Fig. 1C) are two of the best-known predicted Coulomb phases, neither has a good 58 Theoretical investigations of the pyrochlore Heisenberg experimental model system. antiferromagnet highlight the generalities of Coulomb phases [11,12,13,14,15]. The spin 59 60 correlation function has the power-law behavior generic to all Coulomb phases, implying that there should be `pinch-points' in the static structure factor [3,12,16,17]. 61 Creation of 62 topological defects in the flux fields (monopoles, though in this case not discretized as they are in spin ice [18,19,20]) modifies the magnetic charge density locally, and relaxation of this 63 64 disturbance leads to a monopole current. The associated spin relaxation rate depends only on 65 temperature (i.e. it exhibits E/T scaling) and occurs at many wave vectors, including the pinch 66 points, as shown by the theoretical spin relaxation function $S(\mathbf{Q},t)/S(\mathbf{Q},0)$ [14,15] illustrated in 67 Fig. 2A. On nodal lines connecting the pinch points, different behaviors are found. In a cut 68 through the simulated $S(\mathbf{Q}, \omega)$ along a nodal line, as shown in Fig. 2B [15], the relaxational 69 response at the zone centre gives way to spin diffusion at small wave vectors where fluctuations 70 conserve the local magnetization; fast, quasi-dispersive dynamics due to remnant spin waves 71 appear further along the nodal lines. It was recently suggested that the lattice dynamics of 72 solids with correlated disorder (including charge ices) may be of interest for functional 73 properties such as thermoelectricity [21], but the vibrational analogs of general Coulomb phase 74 dynamics have not been investigated. Moreover, in dense spin systems with mixed cations 75 such as Y₂CrSbO₇ or YbMgGaO₄, site and bond disorder clearly have important consequences for the magnetic properties [22,23,24]. These materials are suggested to have random disorder, 76 77 but although correlated disorder may be difficult to detect, it is highly probable because the 78 long-range nature of the Coulomb interaction makes it divergent in the presence of random 79 disorder. Further investigations of such materials may well emphasize the importance of 80 understanding spin systems that live on structures with correlated disorder.
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82 In this work, we use x-ray and neutron scattering to investigate correlations and dynamics in 83 CsNiCrF₆, in which Ni²⁺ and Cr³⁺ jointly occupy a pyrochlore lattice. CsNiCrF₆ was suggested 84 to be a pyrochlore Heisenberg antiferromagnet [25,26] before the concept of a Coulomb phase 85 emerged. However, even in the simplest description [27], Ni-Ni, Ni-Cr and Cr-Cr 86 superexchange constants form a set of different values, breaking the local degeneracy of the 87 true pyrochlore Heisenberg antiferromagnet that is crucial to the formation of a Coulomb 88 phase. However, with two ions of different charge on the pyrochlore lattice, CsNiCrF₆ is a 89 candidate charge ice and later we show how a magnetic Coulomb phase is inherited from the 90 charge ice structural correlations.

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92 The charge ice structure can be addressed in two ways: the average crystal structure can be 93 studied by conventional diffraction experiments, and departures from the average due to

94 (correlated) disorder can be investigated by measuring structural diffuse scattering [28]. To 95 investigate the average structure, we performed single crystal neutron diffraction and 96 synchrotron powder x-ray diffraction experiments. Refinement of crystallographic models 97 against these data indicate three features: firstly, the average structure is well described by that 98 of the mixed metal fluoride pyrochlores (see Fig. 1D,E), with isotropic displacement 99 parameters [29]; secondly, the thermal displacement factor of the Cs⁺ ion is by far the largest of all the atoms, and is strongly thermally activated; lastly, split site models in which ions 100 101 partially occupy a lower symmetry site around the expected position, as in other pyrochlore 102 materials with structural disorder [30] are not stable or do not improve the refinement. Further details of the refinements are given in the supplementary information [31]. 103

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105 If there is sufficient contrast in neutron scattering length between the two cations, as is the case 106 for Ni ($b_c=10.3$ barn) and Cr ($b_c=3.6$ barn), there should be a direct signature of charge ice 107 correlations in the structural (i.e. nuclear) diffuse neutron scattering. We used polarized neutron scattering to separate structural and magnetic scattering. The structural diffuse 108 109 scattering is shown in Fig. 3A, and its form indicates that non-trivial structural correlations are 110 present. In the experimental data the intensity of the diffuse features generally increases with the magnitude of the wavevector, while a model calculation of a charge ice with only 111 112 substitutional correlations (Fig. 3B) has diffuse features with similar weight across the whole pattern. The calculation has pinch points at (-2, -2, 0) and (0, 0, 2), whose existence and position 113 114 are the defining features of a structural Coulomb phase on the pyrochlore lattice [3]. We can 115 identify one of these pinch points in the experimental data. In Fig. 3A, a triangular region of intensity around (h,h,0) sharpens inward, towards (2,2,0). Cuts through the experimental data 116 (Fig. 3D) confirm that the scattering sharpens toward the (2,2,0) Bragg point, and that this 117 118 feature is therefore a type of pinch point. The very weak intensity of the inner part of the pattern prevents us from concluding on the exact nature of the features around (0,0,2), but the 119 120 intense diffuse scattering pinching in toward (0,0,6) at the limit of our wave vector coverage 121 suggests another pinch point, as expected from the calculation. The pinch points are the 122 essential characteristic of the Coulomb phase, indicating the long-range coherence of the ice 123 rules, their presence in the experimental data shows unambiguously that a structural Coulomb 124 phase is present.

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126 The other, more diffuse, features characterize short-wavelength correlations or local structures 127 that exist within the Coulomb phase, and the distribution of intensity in the experimental pattern compared to the calculation shows that the structure contains contributions beyond 128 129 purely ice-rule obeying cation configurations. When working with single crystal diffuse 130 scattering data with relatively limited wave vector access, numerical modelling in direct space 131 is most convenient [32, 33, 34, 35], and we propose a simple model that explains our structural 132 observations reasonably well. From the point of view of the bond valence sum [36], the average structure is an efficient compromise: no cation has a fully favorable bond valence sum, 133 134 but all are close to their optimal values. If ice rule cation correlations exist amongst Ni²⁺ and 135 Cr^{3+} , we can make a local distortion throughout the structure (Fig. 1D) that brings the bond 136 valence sum of both cations to their optimal values. The Ni/Cr cations have octahedral 137 coordination, sharing four F with cations of the other type and two with cations of the same type. We assume only that any F^{-} shared between a Ni²⁺-Cr³⁺ pair is displaced (always by the 138 139 same amount) toward the Cr³⁺, sufficient to simultaneously optimize the bond valence sum of 140 all Ni²⁺ and Cr³⁺ ions. The displacement is small enough to lie within the radius of the isotropic 141 displacement parameter, consistent with the absence of a split site in structural refinements or 142 Fourier difference maps. Then, if we search for the locus of sites optimizing the bond valence 143 sum for each Cs⁺, we find it takes the form of one or more connected loops on the surface of a sphere, with the average over all Cs^+ sites making a hollow spherical shell. As in other such cases, the radius of the shell is larger than the displacement parameter of the atom in the average

146 structure [37].

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148 The introduction of the cation correlations and local distortion of the octahedra has a strong 149 effect on the calculated structural diffuse scattering - the local distortions redistribute the 150 weight of the pure configurational disorder pattern towards larger wave vector, suppressing the 151 part in the center of the pattern (Fig. 3C) but leaving the pinch point at (2,2,0) described above. 152 Furthermore, the combination of cation correlations and local anion displacements modifies the topology of the available Cs⁺ positions from discrete pockets in the average structure (i.e. 153 154 a split site), to continuous displacements on the surface of a sphere, which on average resembles 155 a large, isotropic displacement parameter as observed. If the framework cations are random, the local distortions still improve the bond valence sums, and still have the same effect on the 156 157 Cs⁺ displacement topology, but the diffuse scattering is a featureless response from the random 158 cations convoluted with weighting to larger wave vector due to the displacements.

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160 Every tetrahedron has four edges with a displacement and two without, a condition that can 161 readily be represented by an ice rule on a single tetrahedron, but the relationship between ice 162 rule and fluxes requires a sign change in `up' and `down' tetrahedra, since anions are displaced 163 toward a cation in both of the tetrahedra of which the cation is a member. An alternative 164 identification of the Coulomb phase in the displacements can be made: the edges of the 165 tetrahedra with a non-displaced F- anion (or matching cations at each end) select two of the 166 links of the diamond lattice that terminate at the diamond lattice point at the center of the tetrahedron (Fig. 1D); each cation is a member of another such pair on adjacent tetrahedra, 167 similarly selecting two diamond lattice links on these tetrahedra (Fig. 1D, E); tracing the 168 169 network of diamond lattice links selected in this way reveals a fully packed loop model. Every link of the diamond lattice is a member of a loop and each diamond lattice point is visited by 170 171 precisely two loops; a given loop is populated only by one type of cation. The same fully 172 packed loop model is obtained by coloring, in two colors, the links of the diamond lattice that 173 host in or out spins in a spin ice [38]. Fully packed loop models represent an intriguing type 174 of correlated disorder, which has so far found experimental relevance only in two dimensions [39,40]. In CsNiCrF₆, this displacement ice must play an important role in the exchange 175 176 interactions and their disorder. The ideal pyrochlore Heisenberg antiferromagnet has a single 177 exchange parameter, but here we can expect three, one each for Ni-Ni, Ni-Cr and Cr-Cr bonds. 178 The required theory will not be one of random disorder of exchange strengths, but of a strongly 179 correlated spatial distribution of exchanges, whose structure is encoded by the fully packed 180 loop model [27].

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182 Our model does not reproduce every aspect of the structural diffuse scattering pattern, but the form of the experimental data and improvements made over the simple charge ice calculation 183 suggest the basic ingredients – a charge ice of Ni²⁺ and Cr³⁺, displacements of the F⁻ anions that 184 inherit Coulomb phase correlations from the underlying charge ice, and Cs⁺ ions disordered 185 186 over the surface of a spherical shell around the cage centers – are correct. However, the model 187 is very simple: the ice rules are assumed to operate perfectly (some additional consideration of 188 this point can be found in the supplementary information [31]), with complete degeneracy 189 amongst all microstates (which is known not to be the case in spin ice); the same F-190 displacement is made throughout the structure, irrespective of the symmetry of the local 191 distortion - $O_h \rightarrow D_{4h}$ and $O_h \rightarrow C_{2\nu}$ may both occur depending on the configuration of 192 displaceable F⁻ ions around each Cr³⁺ and may in reality be differently favorable; and the Cs⁺ 193 ions are completely decoupled from the framework.

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195 From the point of view of magnetism, the Curie-Weiss temperature (-70 K) suggests that in 196 CsNiCrF₆, the magnetic moments of two sizes (Ni²⁺ has S=1, Cr³⁺ has S=3/2) distributed according to the ice rules, interact antiferromagnetically on average. Separation of the 197 198 magnetic diffuse scattering by polarized neutron scattering shows that the spins also form a 199 Coulomb phase. The static magnetic correlations exposed in a measurement of $S(\mathbf{Q})$ contain 200 pinch points, and a distribution of intensity similar to the pyrochlore Heisenberg 201 antiferromagnet, as shown in Fig. 4A and B. The existence of pyrochlore Heisenberg antiferromagnet-like correlations in a system with two different interacting ions (distributed 202 203 according to the ice rules/fully packed loop model) has been justified by numerical simulations 204 for an extensive range of interaction parameters (see regime IV in Ref. [27]). It is clear from 205 our results that CsNiCrF₆ falls within this range. As discussed in Ref. [27], the Coulomb phase 206 correlations of the spin system are here inherited from the underlying structural Coulomb phase 207 and do not reflect the degeneracy of the Heisenberg manifold.

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209 This conclusion is supported by a study of the magnetic dynamics using inelastic neutron 210 scattering. Previous studies have shown that there is considerable spectral weight at an energy 211 transfer of ~0.5 meV [26], and in Fig. 5A we show a map of the intensity at constant energy 212 transfer of 0.5 meV. This measurement shows that the spin fluctuations have a structure factor very similar to the static correlations of the Coulomb phase, implying degenerate fluctuations 213 214 of both long and short wavelength. Constant wave vector scans were used to further investigate 215 the form of $S(\mathbf{Q},\omega)$. The locations of these scans are summarized in Fig. 2A, they probe the 216 nodal lines (h,h,-h) and (0,0,l), including pinch points at (1,1,-1) and (0,0,-2), and generic points 217 on the intense structure factor of Fig. 5A. In Fig. 5B, we show scans measured at the pinch points and generic points. We have converted $S(\mathbf{Q},\omega)$ to $\chi''(\mathbf{Q},\omega)$ in order to simplify model-218 219 free interpretations. The data from all these points can be fitted by gapless quasi-elastic 220 Lorentzians with the same width, indicating that the relaxation rate is independent of wave 221 vector (but the intensity varies with position, as expected from Fig. 5A). In Fig. 5C we show 222 a series of scans along a nodal line from a pinch point at (1,1,-1) toward the adjoining regions 223 with no scattering in Fig. 5A (i.e. (0,0,0) or (2,2,-2)). Here we see that the gapless behavior at 224 the pinch point is preserved at small wave vector relative to the pinch point (e.g. (1.1,1.1,-1.1), 225 but a broad peak with a distinct dispersion appears beyond this. Beyond (1.5, 1.5, -1.5), the peak 226 becomes very weak and much broader, dispersing back downward as far as it can be followed. 227 The temperature dependence of the width of the susceptibility at two generic wavevectors is 228 linear with the temperature, as shown in Fig. 5D, with a finite response at the lowest 229 The width of the dispersive feature has essentially the same temperature temperature. 230 dependence. In Fig. 5E we summarize our data for nodal lines, showing the sinusoidal 231 dispersion of the fast dynamics.

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233 CsNiCrF₆ affords the possibility of studying the dynamics of a magnetic Coulomb phase with 234 full wave vector and frequency information. Interestingly, many features of these dynamics 235 are similar to those of the classical spin liquid phase of the pyrochlore Heisenberg 236 antiferromagnet. In particular, monopole current density is signified by the wave vector-237 independent relaxation rate with linear temperature at generic wave vectors [14]. Along the 238 nodal lines, the response at and close to the pinch point is gapless. This is consistent with the 239 presence of relaxational dynamics at the pinch point and diffusional dynamics at small wave 240 vector but more extensive measurements would be required to confirm the expected data 241 collapse for spin diffusion. We clearly see a feature that disperses up from the pinch point. 242 The dispersion on the nodal lines is periodic with respect to the pinch points (and identical for 243 the two nodal lines). When compared qualitatively with the prediction of Fig. 2B [14, 15], the observation of a quasi-linear or sinusoidal dispersion up to the maximum, followed by a collapse into a very broad, weak and downward dispersing intensity is as expected. This dispersive feature represents a fast contribution to the dynamics that can be thought of as a spin wave fluctuation about the local ground state. It is broad in both wave vector and energy since the ground state is not ordered [14].

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250 It may seem remarkable that the delicate-sounding predictions of the pyrochlore Heisenberg 251 antiferromagnet are robust to the strongly correlated structural disorder that we have also 252 described. However, the fully packed ion loops of a charge ice still give rise to a magnetic 253 Coulomb phase of antiferromagnetic character [27], and conserved or topological quantities 254 are closely related to those of the Heisenberg model, leading to similar emergent physics. The 255 expected clear departure from the classical pyrochlore Heisenberg antiferromagnet behavior 256 is, however, manifest in the temperature dependence of the relaxation rate at the generic 257 wavevectors. This quantity is predicted to be simply proportional to the temperature and vanish 258 as $T \rightarrow 0$ [14,15], but we find a weaker temperature dependence and residual response at the 259 lowest temperature.

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Below T=2.3 K magnetization measurements on CsNiCrF₆ exhibit a history dependence that is 261 262 reminiscent of canonical spin glasses [25]. A slowing component of dynamics that coexists 263 with the processes we measure here, and which falls into the time window of a magnetization 264 experiment at this temperature seems by no means inconsistent with Ref. [25], but the 265 significant inelastic spectral weight that we observe both above and far below this temperature precludes that CsNiCrF₆ is a canonical spin glass [25,26]. Given the relatively small spin 266 267 sizes, the lowest temperature response may indicate that a quantum spin liquid [41] replaces 268 the classical one.

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270 We have shown that CsNiCrF₆ supports multiple Coulomb phases – a charge ice and associated 271 displacement ice, and a magnetic Coulomb phase with antiferromagnetic character. Despite 272 being composed of fully-packed loops of distinct spins, the magnetic Coulomb phase shows 273 dynamical signatures of the pyrochlore Heisenberg antiferromagnet, a classical spin liquid of 274 much higher local symmetry. Our results show how two systems that look completely different 275 from a structural chemistry perspective, may nevertheless share many of the same physical 276 properties, the key connection being the robust local gauge symmetry of the Coulomb phase 277 and its associated conservation laws.

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279 Predictions of wave vector-dependent dynamics of a classical antiferromagnetic Coulomb 280 phase have not previously been examined experimentally (although we note very recent work on a possible quantum analogue, NaCaNi₂F₇ [42]). The consequences of ice-rules disorder in 281 282 the vibrational dynamics (of the framework) have not been examined experimentally or 283 theoretically, but the interplay of framework and caged-atom dynamics are well known to 284 contribute to advantageous thermoelectric properties in skutterudite and clathrate compounds 285 such as CeFe₄Sb₁₂ [43], and superconductivity in the analogously structured osmate 286 pyrochlores such as KOs₂O₆ [44]. The chemical flexibility of the fluoride pyrochlore crystal 287 structure (which includes the possibilities of significantly varying the magnetic properties with 288 numerous combinations of transition metal cations, replacing the alkali metal cation with a 289 guest of quite different character such as NH₄⁺, or extracting it, as in FeF₃, and of exploring 290 charge ordered versions using compositions such as AFe_2F_6 [45,46], perhaps with a view to 291 metallization) suggests that these compounds are of considerable interest for exploration of 292 both unconventional spin dynamics, and new functionalities based on correlated disorder.



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294 Fig. 1. Illustration of Coulomb phase construction/mappings and structural features of 295 CsNiCrF₆. A: The pyrochlore lattice (medial) consists of corner-sharing tetrahedra, on which 296 a Coulomb phase is established when the local degrees of freedom can be represented by a 297 non-divergent field derived from lattice fluxes on the diamond lattice (parent) [3]. A non-298 divergent configuration of flux variables (blue arrows) around a diamond lattice point is 299 equivalent to the ice rule for spins in a spin ice (two-in-two-out). Antiferromagnetic pseudospins are related to the flux variables and configured two-up-two-down in a Coulomb phase 300 ground state (green and orange arrows). B: Two cations occupying the pyrochlore lattice form 301 a charge ice, with cation configurations directly related to the pseudo-spins (green and orange 302 spheres). C: Vector spins coupled antiferromagnetically on the pyrochlore lattice (i.e. a 303 304 pyrochlore Heisenberg antiferromagnet) must satisfy the condition that the total magnetization 305 of every tetrahedron is zero (black arrows), which can be ensured if the $S_{x,y,z}$ vector components are each represented by a family of pseudospins that obey the ice rule (blue, green and orange 306 arrows). D: In the structure of CsNiCrF₆, F⁻ anions (small blue spheres) are coordinated to two 307 cations (orange spheres are Cr³⁺, green spheres are Ni²⁺). The bond valence sum requirements 308 309 of the cations can be satisfied by displacing anions shared by a pair of unlike cations towards 310 the cation with larger charge (Cr^{3+}) . Anions shared between a pair of like cations are not 311 displaced. The edges with non-displaced anions form a fully packed loop model (described in 312 text) on the diamond lattice (green and orange lines). E: The structure of CsNiCrF₆ is 313 composed of the pyrochlore lattice of cations, coordinated by octahedra of F⁻ ions. The 314 octahedra share each of their vertices with another octahedron, forming tetrahedra of tilted 315 octahedra. (Extended sections of the fully packed loop model are shown and the octahedra are 316 drawn undistorted.) F: The structure contains large cages in which the Cs⁺ ions reside. The 317 cage coordination is by 18 F⁻ ions, forming an octahedron that is truncated and capped with 318 rectangular pyramids.

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Fig. 2. Illustration of theoretical predictions of the dynamical response of the pyrochlore Heisenberg antiferromagnet. The relaxation function $S(\mathbf{Q},t)/S(\mathbf{Q},0)$ (from [15]) has large regions of reciprocal space where identical relaxation times are expected. The normalization highlights that the scattering function at these wavevectors will have the same width, even if their structure factor is different. The nodal lines, where different dynamical regimes are found, can be clearly seen. The extent (curved lines) and positions (dots) of our mapping and scanning measurements are also shown. B: A sketch of the spectrum along a nodal line (i.e. (h,h,-h)) (from [15]) shows the crossovers from gapless relaxational dynamics at the pinch point $(\mathbf{k}_{nodal}=0)$, to gapless diffusional dynamics at small $|\mathbf{k}|$, to quasi-dispersive fast dynamics toward $|\mathbf{k}|=0.5$, with broad and weak signal for $|\mathbf{k}|>0.5$.





Fig. 3. Structural diffuse scattering in CsNiCrF₆. A: The structural diffuse neutron scattering 343 344 measured at 1.5 K (note that the colour scale starts at 0.01 (arbitrary units) to highlight the weak diffuse scattering around (1,1,1) and (0,0,2), there is a significant wavevector 345 346 independent diffuse contribution which may arise from isotope incoherent scattering, 347 uncorrelated Cs⁺ ions, and/or ice rule defects). B: Predicted diffuse scattering of a charge ice 348 with substitutional correlations only. C: Diffuse scattering calculated for a model with charge 349 ice cation correlations and associated bond valence sum-restoring F⁻ and Cs⁺ displacements. In 350 B and C the calculated intensity is multiplied by a scale factor and a wavevector independent 351 contribution added. D: Cuts through the experimental data (positions indicated by lines on A) 352 show that the feature around (h,h,0) sharpens inward toward the (2,2,0) position, and is, 353 therefore, the pinch point visible at that position in B and C. Error bars in D are obtained by 354 standard propagation of the Poisson counting statistics of all the measurements (sample, 355 background, calibrants) required to extract the structural scattering cross section.





Fig. 4. Magnetic diffuse scattering in CsNiCrF₆. A: The experimental magnetic diffuse neutron scattering measured at 1.5 K. B: Model calculation of the structure factor of the pyrochlore Heisenberg antiferromagnet (including the average magnetic form factor for Ni²⁺ and Cr³⁺) [16]. The calculation is for T=0 and incorporates no structural disorder or modulation of the magnetic moment size by site, so its pinch points are extremely sharp and better defined than

362 those in the experimental data. Isointensity contours from the calculation are superimposed on 363 the experimental data as guides to the eye.



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365 Fig. 5. Wave vector and temperature dependence of magnetic dynamics in CsNiCrF₆. A: A map of $S(\mathbf{Q},\omega)$ at a constant energy transfer of 0.5 meV shows a structure factor very similar 366 to the static correlations shown in Fig. 4, but with broadened pinch points. B: Constant wave 367 vector scans at pinch points and points on the structure factor of panel A where relaxational 368 dynamics are expected (see Fig. 2) all have the same width (small numbers adjacent to a data 369 370 series indicate an arbitrary offset that has been applied to help distinguish them, the real 371 intensity of these series is essentially identical, lines are fits to a gapless quasielastic Lorentizan line shape). C: Constant wave vector scans along the nodal line (h,h,-h) show a gapless 372 373 spectrum at small **k**, followed by an upward dispersive peak for 0.2<**k**<0.5, followed by 374 broad, weak signals of decreasing intensity (lines are fits to a damped harmonic oscillator line 375 shape with pole at finite energy). D: The width of the relaxational dynamics signal is linear in 376 T, with a significant intercept at low temperature. The width of the dispersive peak at (1.5, 1.5, -5)1.5) has an essentially identical temperature dependence. E: Summary of the nodal line 377 378 dynamics, showing gapless behavior close to the pinch point, replaced by dispersive fast 379 dynamics and weak broad scattering. The points and line indicate the dispersion of the 380 susceptibility peak along both nodal lines and in both directions from the pinch point, folded into reduced units. The color map is an interpolation of the susceptibility measured in the 381 382 constant wave vector scans (along (h,h,-h)). Error bars in B and C are derived from Poisson counting statistics of the measurement with scaling from $S(\mathbf{Q},\omega)$ to $\chi''(\mathbf{Q},\omega)$. Error bars in D 383 384 and E are the uncertainties of the least squares fitted parameters.

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387 Materials and Methods [References 47-56]:

388 Our main sample is a single crystal of $CsNiCrF_6$, used and described in [25,26]. It is roughly 389 cuboidal, with dimensions $\approx 6x8x12$ mm. It was aligned by neutron Laue diffraction using 390 Orient Express at the Institut Laue Langevin (ILL) (Grenoble, France) and mounted according 391 to the intended experiment: glued to an aluminum finger with Quikfill (epoxy) for inelastic 392 neutron scattering experiments at T>1.5 K; glued to a copper finger with Stycast (epoxy) for 393 inelastic neutron scattering experiments at T < 1 K; wrapped in aluminum foil that was clamped 394 to an aluminum finger for polarized diffuse neutron scattering experiments. Small pieces of 395 dimension $\approx 0.5 \times 1 \times 2$ mm that broke off the surface of the sample during ungluing were 396 preserved and used for single crystal neutron diffraction experiments (glued to aluminum pins 397 using Quikfill or GE varnish) or crushed for powder x-ray diffraction. We have a second 398 crystal that has a well-developed octahedral form with edge dimension ≈ 5 mm. A small piece 399 cut from this was also examined by x-ray powder diffraction.

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401 For our synchrotron x-ray powder diffraction experiments, a piece of either crystal was mixed 402 with silicon powder and ground together in an agate pestle and mortar. These mixtures were 403 loaded into 0.3 mm glass capillaries. The silicon serves primarily to disperse small sample 404 volume in the beam, and also provides a convenient calibrant for wavelength and lattice 405 parameters (a_{si}=5.431194 Å at 22.5°C, NIST powder diffraction standard 640c). We used the 406 high-resolution powder diffractometer of the Materials Science Beamline [47] at the Swiss 407 Light Source (SLS) (Paul Scherrer Institut, Villigen, Switzerland) to measure the diffraction 408 pattern of the mixture. The diffractometer was operated in Debye-Scherrer geometry with 409 Mythen microstrip detector and capillary spinner, the wavelength was 0.4959 Å (25 keV). The 20 range extended from 7° to 120° without cryostat, or 7° to 80° with helium flow cryostat 410 installed. We collected diffraction patterns at room temperature (24°C at the SLS) without the 411 412 cryostat for the main sample, and between 5 K and 300 K with the cryostat for both. The 413 powder diffraction data were normalized and reduced by standard routines, then modeled and 414 fitted using the Rietveld method, as implemented in the package FullProf [47].

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416 We carried out a single crystal neutron diffraction experiment using the quasi-Laue 417 diffractometer VIVALDI [49] at the ILL. The sample was mounted in an `Orange' helium 418 cryostat and cooled to $T \approx 2.5$ K. To ensure a full coverage of reciprocal space, the sample was 419 rotated through 180° about the vertical axis, with 1-hour duration exposures recorded every 10°. The data were indexed and integrated using the program LAUEGEN [50], and wavelength 420 421 normalized using LAUENORM [51]. The resulting integrated intensities were used for least 422 squares refinement of crystallographic models using the package ShelXL [52]. A second single 423 crystal neutron diffraction experiment was carried out using the monochromatic diffractometer 424 TRiCS [53] at the Swiss Spallation Neutron Source (SINQ) (Paul Scherrer Institut, Villigen, 425 Switzerland) in 4-circle mode. The sample was attached to the cold finger of a closed cycle 426 cooling machine that was mounted on the Euler cradle. Using the germanium (Ge311) monochromator to provide neutrons of wavelength of 1.172 Å and a single ³He tube detector, 427 428 we measured 2550 reflections, with the temperature T=5 K throughout. Integrated intensities 429 were combined and used for least squares refinement of crystallographic models using the 430 packages Jana2000 and FullProf [54,47].

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432 We measured and separated the magnetic and structural diffuse neutron scattering using the

433 polarized neutron diffuse scattering spectrometer D7 [55] at the ILL. D7 was configured with 434 wavelength λ =3.1 Å and Orange cryostat. The sample was mounted using aluminum foil to

434 wavelength λ =3.1 Å and Orange cryostat. The sample was mounted using aluminum foil to 435 minimize incoherent scattering from glue, two strong Bragg peaks from the tails of the cryostat 436 contaminate the data and these areas are masked in the data analysis. We used the XYZ 437 technique, which requires the measurement of non-spin flip and spin flip intensities for three 438 orthogonal polarization directions, to separate the structural, spin incoherent, and magnetic 439 cross sections. We rotated the crystal about the vertical axis in 0.5° steps, recording three 440 complete rotations with the analyzer/detector banks offset in three non-overlapping positions 441 and 17-18 s count time per channel, plus a further 270° rotation in one of the detector positions 442 with 20 s count time. The sample was maintained at $T \approx 1.8$ K throughout. We measured the 443 background of the empty sample holder for each of the three detector positions with coarse 444 rotation steps (18° or 30°) and 60 s counting times. The background was essentially angle 445 independent so measurements at different rotation angles were combined and duplicated for 446 subtraction from the sample data. Standard vanadium and quartz (silica glass) samples were 447 used for normalization of detector and polarization analyzer efficiencies respectively. The 448 nuclear, magnetic and spin incoherent cross sections were separated [55] and symmetrized by 449 folding into a single quadrant of the scattering plane and then unfolding.

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451 We report data from three inelastic neutron scattering experiments performed using the cold 452 neutron triple axis spectrometer IN14 at the ILL. In the first, the sample was mounted in an Orange cryostat and cooled to $T \approx 1.8$ K. The instrument was configured with PG002 453 454 monochromator and FlatCone analyzer. The FlatCone analyzer has 31 silicon (1,1,1) analyzer 455 crystals, allowing to map the excitation spectrum at a single energy transfer. We recorded a 456 145° rotation of the sample (in 1° steps) with the analyzer bank offset at two positions. A 457 measurement of the incoherent scattering from a vanadium sample was used to normalize the efficiency of the analyzer-detector channels. We performed two experiments in the 458 459 conventional triple axis configuration with PG002 monochromator and analyzer. In one, the 460 sample was cooled using the Orange cryostat, and our main collection of constant-wave vector scans at various reciprocal lattice positions and temperatures was made. In the other, the sample 461 was cooled using a dilution refrigerator insert in the Orange cryostat to reach $T \approx 0.07$ K, and a 462 463 limited collection of scans was made. We estimate the energy resolution at the elastic line to be $\Delta E \approx 0.1$ meV when $k_f = 1.5$ Å⁻¹ and $\Delta E \approx 0.06$ meV when $k_f = 1.3$ Å⁻¹. 464

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466 Our numerical model of the structural diffuse scattering was encoded specifically by ourselves. The average crystal structure is built in a cubic supercell of the pyrochlore unit cell of size 467 468 L=6,12. We establish an ice rule configuration of cations on the pyrochlore lattice by first tiling it with hexagonal loops (supercell sizes of L=6,12 are required to ensure complete 469 470 coverage, such a tiling is illustrated in Ref. [56]). Starting in the lowest layer of the lattice, a 471 hexagon-loop covering is constructed with probabilistic choice between two possible origins. 472 On moving up to the next layer where sites are not covered by hexagons, the layer configuration 473 is again chosen probabilistically. Once every site of the whole lattice is assigned to a hexagon, 474 the covering can be permuted, again probabilistically, so that each possible growth direction of 475 the hexagon coverage is equally represented. Within individual hexagons sites are assigned 476 alternating up/down pseudo-spins, again choosing probabilistically between the two possible 477 configurations. At this point, a specific ice-rule obeying pseudo-spin configuration has been 478 generated, but one which does not have power-law correlations. The full power-law correlation 479 function is obtained by propagating a large number of loop moves, allowing loops of any length Simulation of the magnetic neutron scattering pattern using spin 480 and winding loops. 481 configurations generated in this way will result in the T=0 scattering pattern of the pyrochlore 482 Heisenberg antiferromagnet or spin ice, depending which type of spin is associated with the pseudo-spin configuration. For the structural simulation, pseudospin orientation is converted 483 into cation identity, and we identify those tetrahedron edges with mixed cation pairs and shift 484

- the relevant F⁻ anion towards its neighboring Cr^{3+} cation. Calculation of the bond valence sums at this point reveals universal satisfaction of the requirements of Ni²⁺ and Cr³⁺, but Cs⁺ remains unsatisfied. We search systematically around every cage center for positions that satisfy the Cs⁺ bond valence sum, within a certain tolerance, and with a certain resolution in spherical
- 488 Cs^+ bond valence sum, within a certain tolerance, and with a certain resolution in spherical 489 polar coordinates, and tabulate all these positions. Then we choose positions for each Cs^+
- 490 randomly from its list and calculate the scattering pattern. The calculated data shown in Fig.
- 491 2B were obtained by repeating this procedure 200 times with L=12. The model intensities are
- 492 scaled and a constant background added to compare with the experimental data.
- 493

494 Data availability: The experimental data and their supplementary information, analyses and
 495 computer codes that support the plots within this paper and the findings of this study are
 496 available from the corresponding author upon reasonable request.

497

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- 505 **Author notes:** TF, MJH, SC, MB, PS, and STB carried our inelastic neutron scattering 506 experiments; TF, M.-HL-C, and OZ carried out neutron diffraction experiments; MR and AC 507 carried out x-ray diffraction experiments; TF analyzed all data and made calculations; TF, 508 MJH, and STB wrote paper in collaboration with all other authors.
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