1	Hydrogens and hydrogen-bond networks in macromolecular MicroED data
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13	Abstract
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15	Microcrystal electron diffraction (MicroED) is a powerful technique utilizing electron cryo-
16	microscopy (cryo-EM) for protein structure determination of crystalline samples too small for X-
17	ray crystallography. Electrons interact with the electrostatic potential of the sample, which means
18	that scattered electrons carry informing about the charged state of atoms and can provide strong
19	contrast for visualizing hydrogen atoms. Accurately identifying the positions of hydrogen atoms,
20	and by extension the hydrogen bonding networks, is of importance for drug discovery and
21	electron microscopy can enable such visualization. Using subatomic resolution MicroED data
22	obtained from triclinic hen egg-white lysozyme, we identified hundreds of individual hydrogen
23	atom positions and directly visualize hydrogen bonding interactions and the charged states of
24	residues. Over a third of all hydrogen atoms are identified from strong difference peaks, the most
25	complete view of a macromolecular hydrogen network visualized by electron diffraction to date.
26	These results show that MicroED can provide accurate structural information on hydrogen atoms
27	and non-covalent hydrogen bonding interactions in macromolecules. Furthermore, we find that
28	the hydrogen bond lengths are more accurately described by the inter-nuclei distances than the
29	centers of mass of the corresponding electron clouds. We anticipate that MicroED, coupled with
30	ongoing advances in data collection and refinement, can open further avenues for structural

biology by uncovering and understanding the hydrogen bonding interactions underlying protein

32 33 structure and function.

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Microcrystal electron diffraction (MicroED) has been successful in structure determination of 36 crystalline biological specimens using electron cryo-microscopy (cryo-EM) (Nannenga, Shi, 37 Leslie et al., 2014; Nannenga, Shi, Hattne et al., 2014; Yonekura et al., 2015), including novel 38 structures (Rodriguez et al., 2015; Sawaya et al., 2016; Xu et al., 2019; Clabbers et al., 2021), as 39 40 well as difficult to crystallize membrane proteins in detergents and lipids (Liu & Gonen, 2018; Martynowycz et al., 2020; Martynowycz, Shiriaeva et al., 2021). As electrons interact more 41 42 strongly with matter than X-rays (Henderson, 1995), the crystal volume required for useful diffraction is typically about a million times smaller. Electrons are scattered by the electrostatic 43 44 potential and the strength of scattering depends on the charged state of atoms (Cowley, 1995). The effects of charge distribution are already apparent at moderate to low resolution (Yonekura 45 et al., 2015, 2018), and the charged state of residues in macromolecules has previously been 46 investigated using electron crystallography (Kimura et al., 1997; Mitsuoka et al., 1999; 47 Yonekura et al., 2015). 48

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50 Electrostatic potential maps obtained from electron scattering can provide strong contrast for identifying hydrogen atoms, which has enabled localizing hydrogens in electron diffraction 51 structures of small molecule organics and peptide fragments (Rodriguez et al., 2015; Sawaya et 52 53 al., 2016; Dorset, 1995; Palatinus et al., 2017; Gruene et al., 2018; Jones et al., 2018; Clabbers et al., 2019; Takaba et al., 2021). Identifying the positions of hydrogen atoms and visualizing their 54 resulting hydrogen bonding networks are crucial for understanding protein structure and function 55 such as resolving precise drug or ligand binding interactions (Purdy et al., 2018; Clabbers et al., 56 2020; Martynowycz, Shiriaeva et al., 2021) or elucidating mechanisms for substrate transfer in 57 membrane protein transporters and channels (Gonen et al., 2005; Liu & Gonen, 2018). In single-58 particle cryo-EM imaging, individual hydrogen atom positions were localized from 59 reconstructions of apoferritin at 1.2 Å resolution (Nakane et al., 2020; Maki-Yonekura et al., 60 2021) and for the GABA_A receptor at 1.7 Å resolution (Nakane et al., 2020). Here, hydrogen 61 atoms were identified by omitting them from the model and inspecting the peaks in a calculated 62 $F_0 - F_c$ difference map following refinement in Servalcat based on crystallographic refinement 63 routines implemented in REFMAC5 (Murshudov et al., 2011; Yamashita et al., 2021). Since 64 65 resolution is a local feature in cryo-EM, the accuracy of hydrogen identification varies across the 66 map.

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Visualizing hydrogen atoms in macromolecular crystallography generally requires (sub-) atomic resolution data, and the accuracy of localizing hydrogens varies with local structural flexibility which is reflected by the temperature factors. Typically, crystals of macromolecules are more disordered than peptides or small molecules and have a much higher solvent content. Therefore, in absence of atomic resolution data, identification of hydrogen atoms in macromolecular MicroED structures has remained elusive.

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Identifying hydrogen atoms in MicroED data 76

Recently, we reported the structure of triclinic hen egg-white lysozyme at 0.87 Å resolution 78 using electron-counted MicroED data (Martynowycz, Clabbers, Hattne et al., 2021). MicroED 79 80 data were collected from 16 crystal lamellae and the structure was phased *ab initio* as described previously (Supplementary Fig. 1) (Martynowycz, Clabbers, Hattne et al., 2021). Following 81 82 density modification individual atoms could be resolved at sub-Ångström resolution, enabling automated model building of the entire structure without reference to a previously determined 83 84 homologous model (Martynowycz, Clabbers, Hattne et al., 2021). The improvement in data accuracy and resolution in this study compared to previous efforts was realized by combining 85 focused ion-beam milling to produce approximately 300 nm thin crystalline lamellae ideal for 86 cryo-EM at 300 kV (Martynowycz, Clabbers, Unge et al., 2021), and collecting data in electron-87 counting mode at a significantly reduced exposure of only 0.64 e⁻.Å⁻² per crystal dataset 88 (Martynowycz, Clabbers, Hattne et al., 2021). A low exposure rate is required for electron 89 90 counting as it ensures that the rate of scattered electrons remains within the linear range of the 91 camera. Lowering the total exposure also reduces the effects of radiation damage that can affect the structural integrity of the protein and the ability to localize hydrogen atoms (Hattne et al., 92 93 2018; Leapman & Sun, 1995).

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We set out to further refine the *ab initio* model resulting from automated building against 95 the subatomic resolution MicroED data to closely examine individual hydrogen atom positions. 96 97 First, the structural model was refined using electron scattering factors, isotropic atomic displacement parameters, and the default riding hydrogen model in REFMAC5 (Murshudov et 98 al., 2011). Twelve alternate side-chain conformations were modeled upon visual inspection using 99 100 Coot (Emsley et al., 2010), and their occupancies were refined. The model was then refined 101 using anisotropic B-factors until convergence (Supplementary Table 1). A crystallographic $mF_0 - DF_c$ difference map was calculated using a model without hydrogen atoms (Yamashita et 102 al., 2021). Peaks in the difference hydrogen omit map at $\geq 2.0\sigma$ were then identified using 103 PEAKMAX (Winn et al., 2011), and those within 0.5 Å distance from any idealized riding 104 105 position were identified as potential hydrogen atoms. In this manner, we located 376 out of 1067 possible hydrogen atoms corresponding to about 35% of the entire structure. Lowering the 106 threshold to 1.0σ revealed a total of 562 hydrogen atom positions, approximately 53%. At 107 contour levels below 2.0σ , the difference map is nosier, increasing the chance of false positives 108 109 and making it more challenging to unambiguously identify peaks as hydrogen atoms. 110 Nevertheless, these results are the most complete hydrogen bonding network visualized to date by macromolecular MicroED (Table 1). 111

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114 Visualizing hydrogens and hydrogen bonding networks

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Overall, the protein main chain is expected to be more rigid than the side chains; we 116 consequently expect more hydrogen atoms to be found in the backbone than in the protein side 117 chains. At the 2.0 σ threshold, we identified 61 out of 141 possible C α -H hydrogens and 76 out 118 of 127 peptide N-H hydrogen bonds corresponding to approximately 43 and 60% of the entire 119 120 structure, respectively (Table 1, Supplementary Tables 2, 3, and 8). The backbone hydrogen atoms are structurally important and can be involved in forming and stabilizing secondary 121 122 structural elements via non-covalent hydrogen-bonding interactions. For example, the structure of lysozyme has two short antiparallel β -strands and we could identify three strong difference 123 124 peaks at $>3.0\sigma$ indicating the positions of those hydrogen atoms involved in hydrogen bonding interactions (Fig. 1a, Supplementary Video 1). The average N-H distance in the β -strands is 125 1.14(26) Å distance, and the distance between the amide group hydrogen donor and carbonyl 126 acceptor is 2.76(9) Å (Table 2). Interestingly, whereas the Asp52 and Gly54 N-H distances are 127 close to the idealized positions, the difference peak for the Asn44 N-H is located at an almost 128 equal distance shared between the donor and Asp52 carbonyl acceptor (Fig. 1a, Table 2). The 129 130 structure of lysozyme is further composed of several short helices, and we could identify a total 131 of 15 hydrogen bonding interactions in the three standard α -helices (Table 2). For example, in the longest 12-residue α-helix we identified 6 out of 10 possible hydrogen bonds based on strong 132 difference peaks at >2.7 σ (Fig. 1b, Supplementary Video 2). The average hydrogen atom peptide 133 N-H distance for the α -helices is 0.97(14) Å with an average distance between donor and 134 acceptor of 2.84(13) Å (Table 2). 135

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137 Higher flexibility and alternate conformations can affect localizing hydrogen atoms in the side chains. Nevertheless, we could successfully localize side-chain hydrogen atoms in the data 138 and identify several hydrogen-bonding interactions between side-chain atoms (Fig. 2, 139 140 Supplementary Table 2). For example, a difference peak at 2.4σ can be resolved between His15-141 NE2 and Thr89-OG1 indicating a possible shared hydrogen bond between both side chains (Fig. 2a). As expected at pH 4.5, the data shows the solvent-exposed histidine to be protonated at 142 ND1, although the hydrogen distance and angle are different from idealized geometry (Fig. 2a). 143 Another example of hydrogen bonding interactions is illustrated for Tyr53-OH acting as a 144 hydrogen donor to Asp66-OD1 with a strong difference peak at 3.4σ (Fig. 2b, Supplementary 145 146 Table 2).

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In single-particle cryo-EM, it was previously observed that acidic side chains were poorly resolved at moderate to low resolution owing to radiation damage and due to the rapid fall off of the electron scattering factors for negatively charged atoms at lower scattering angles (Yonekura *et al.*, 2015, 2018; Maki-Yonekura *et al.*, 2021). In the MicroED data, the acidic aspartate and glutamate residues and their negatively charged side-chain carboxyl groups are generally well resolved (Fig. 2c). Additionally, clear difference peaks at >2.3 σ were identified in the data for the amide side-chain nitrogen for asparagine and glutamine residues, making it possible to
clearly distinguish between the nitrogen and oxygen atoms of the side-chain amide group (Fig.
2c).

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158 Difference peaks were also identified for several water molecules that are involved in hydrogen bonding interactions with the protein backbone and side chains (Fig. 2d, 159 Supplementary Video 3). Such hydrogen bonding networks can act as long-range proton transfer 160 wires. For example, a water molecule is coordinated with the adjacent Ser91, Leu56, and Tyr53 161 residues and shows two strong difference peaks at $\geq 2.7\sigma$ (Fig. 2d). Two additional water 162 molecules show hydrogen atom peaks at $\geq 2.2\sigma$ and are involved in hydrogen bonding 163 interactions with each other and residues of the neighboring protein backbone (Fig. 2d). The O-H 164 hydrogen bond lengths and angles of the water molecules are reasonably close to ideal values, 165 except for one O-H distance for w1001 which is significantly shorter at 0.64 Å. The hydrogen 166 167 bond distance between the w1001-O proton donor and the Tyr53-O proton acceptor is however close to ideal values at 2.75 Å. 168

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171 Hydrogen bond distances

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173 The sheer numbers of hydrogens visualized in this study allow us to measure and report hydrogen bond distances in a way previously not possible in cryoEM (Supplementary Tables 2-174 10; Figure 3). Electrons are scattered by the potential field generated from electron clouds and 175 the nuclei; similar to neutron diffraction. The peaks in an electrostatic potential map are therefore 176 177 expected to reflect the inter-nuclei distances more than distances between centers of mass of electron clouds as observed in X-ray diffraction. We refined the structure using the default riding 178 hydrogen model based on hydrogen distances between the electron cloud centroids using 179 180 restraints derived from X-ray scattering. We analyzed the identified hydrogen atom difference 181 peaks in the data at $\geq 2.0\sigma$ and calculated the average distance for each of the hydrogen bond types (Table 1, Figure 3, Supplementary Tables 2–10). The number of observations for some 182 bond types is insufficient for a rigorous statistical analysis. We do however find an average Ca-183 H distance for the main chain of 1.11(13) Å for 61 hydrogen bonds, compared to idealized values 184 of 0.98 and 1.10 Å for X-ray and neutron diffraction, respectively (Table 1, Figure 3, 185 Supplementary Table 3). The average distance for all N-H bonds is 1.03(16) Å for 83 186 observations, compared to idealized values of 0.98 and 1.10 Å for X-ray and neutron diffraction, 187 respectively (Supplementary Tables 2 and 8). Interestingly, the distances for the amide N-H 188 bonds that are involved in hydrogen bonding interactions with neighboring residues are slightly 189 190 longer compared to those that are not involved in such electrostatic interactions (Table 1, Figure 191 3).

193 These results suggest an elongation of the hydrogen bond lengths compared to the electron cloud centroid distances assumed in the riding hydrogen mode, although the number of 194 observations for each type is rather limited and the standard deviations from the mean value are 195 quite large (Table 1, Figure 3). Nevertheless, we find an overall trend that the C α -H and N-H 196 197 bond lengths are closer inter-nuclei distances (Gruene et al., 2014; Williams et al., 2018). This observation agrees with previous electron diffraction and imaging experiments that show an 198 apparent elongation of the hydrogen bond lengths compared to X-ray diffraction (Clabbers et al., 199 2019; Takaba et al., 2021; Nakane et al., 2020; Maki-Yonekura et al., 2021). Refinement of 200 structural models derived from electron scattering would therefore benefit from more appropriate 201 202 restraints specific for electrons, including a more accurate riding hydrogen model, as well as taking the electrostatic potential of the crystal into account (Yonekura et al., 2015, 2018; 203 Murshudov et al., 2011; Yamashita et al., 2021; Gruene et al., 2014; Williams et al., 2018). 204

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207 Conclusions

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209 The results demonstrate that hydrogen atom positions can be accurately identified in 210 macromolecular MicroED data. As with X-ray crystallography, this will typically require atomic resolution data or better (Walsh et al., 1998; Howard et al., 2004; Wang et al., 2007; Eriksson et 211 al., 2013; Ogata et al., 2015). In comparison, the structure of triclinic lysozyme was determined 212 previously using X-ray diffraction at 120 K and room temperature to 0.93 and 0.95 Å resolution. 213 respectively (Walsh et al., 1998). The single-crystal low-temperature structure is of high quality 214 and generally has more clearly visible hydrogen atoms than the room temperature model merged 215 216 from three crystal datasets. Difference maps contoured at 1.9σ visualize hydrogen atoms in residues within the better-defined regions of the structure, and at 1.8σ contour level, 77 out of 217 127 peptide N-H atoms (61%) are identified (Walsh et al., 1998). The number of hydrogen atoms 218 219 localized in the low-temperature structure is similar to the MicroED structure at comparable 220 resolution, even though the intensity and model statistics are worse (Supplementary Table 1, Supplementary Fig. 1) (Martynowycz, Clabbers, Hattne et al., 2021). The higher level of 221 inaccuracy in the MicroED data can in part be attributed to non-isomorphism from merging of 16 222 crystal datasets and lower completeness in the highest resolution shells (Supplementary Fig. 1). 223 Additional factors that contribute to the errors are inelastic scattering and inaccurate modeling of 224 the electron form factors and the electrostatic potential in structure refinement (Yonekura *et al.*, 225 226 2015, 2018; Murshudov et al., 2011; Yamashita et al., 2021). Compared to X-ray diffraction, electrons are expected to provide better contrast for identifying hydrogen atoms at a similar 227 resolution as the scattering factors fall off less steeply with decreasing atomic number. The 228 lighter hydrogen atoms are therefore expected to be better resolved next to the heavier atoms, 229 230 which might explain why we can identify many hydrogen atoms even though the MicroED data 231 appear noisier. This is further supported by a comparison between apoferritin models from X-ray crystallography and single-particle cryo-EM showed that hydrogen atoms were visibly more 232

clearly in the latter (Yamashita *et al.*, 2021). More recently, a significantly higher resolution structure of triclinic lysozyme was solved *ab initio* at 0.65 Å by X-ray diffraction (Wang *et al.*, 2007). At this resolution, approximately 31% of all hydrogen atoms in main and side chains could be identified at 3.0σ or higher. We would anticipate major improvements in hydrogen atom localization in MicroED data upon further increasing the resolution.

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239 Previously, hydrogen atoms were successfully identified in protein complexes by singleparticle cryo-EM. In comparison to the results presented here, these studies reported that about 240 70% of the expected number of hydrogen atoms could be identified above a threshold level of 241 242 2.0 sing hydrogen-only omit maps from atomic resolution reconstructions of apoferritin (Yamashita et al., 2021; Maki-Yonekura et al., 2021). Remarkably, about 17% of possible 243 hydrogen atoms could be identified from data as low as 1.84 Å resolution (Yamashita et al., 244 245 2021). In imaging, the phase information is retained and during reconstruction, images are 246 filtered to remove noise and to select a specific conformational state. The resolution is therefore 247 a local feature of the map whereas the *B*-factor is a global parameter applied in map sharpening or blurring. This is unlike a crystallographic map, where the resolution is a global feature of the 248 249 entire dataset and structural flexibility or disorder is modeled locally using alternate 250 conformation and per atom refined *B*-factors. In crystallography, resolving detailed features such 251 as hydrogen atoms is affected by local disorder. In the MicroED structure, twelve residues are 252 modeled with alternate side-chain conformations at low occupancy, making it more challenging to identify hydrogen at these positions. The mean *B*-factor over all atoms in the model is 11.98 253 $Å^2$, and the majority of outliers are on the outside of the protein facing the solvent. Lower 254 255 completeness in the higher resolution shells and non-isomorphism from merging data of multiple crystals could both contribute to increased *B*-factors. Especially the last two C-terminal residues 256 have high temperature factors, these were also poorly resolved in the high-resolution X-ray 257 diffraction structure (Walsh et al., 1998). Indeed, most hydrogens can be identified within the 258 259 more stable core of the protein relative to the residues on the outside facing the solvent having higher flexibility and *B*-factors. 260

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In all, 377/1067 (35%) hydrogen atoms could be located at $\geq 2.0\sigma$ and we illustrate 262 263 several examples of well resolved hydrogen atom positions and hydrogen bonding interactions between protein residues and solvent molecules. This is the most complete hydrogen network 264 map for macromolecular MicroED data to date, and these results provide a glimpse of the 265 information that can be obtained by electron scattering, opening up new avenues for further 266 experiments investigating hydrogen bonding networks in protein structures. At the current stage, 267 the difference map becomes increasingly noisy at contour levels below 2.0σ , making it more 268 269 challenging to unambiguously identify hydrogen peaks. Future efforts that can enhance the localization of hydrogen atoms should be focused on improving data accuracy and resolution 270 271 even further. Energy filtration can improve data quality by discarding inelastically scattered 272 electrons, improving the detection of weak peaks at high resolution and at the lower scattering 273 angles that are shaded by the direct beam (Yonekura et al., 2015, 2019). It would also mean the 274 exposure could be lowered even further without losing the weak signal from high-resolution 275 reflections to the noise of the background. Energy filtering does not exclude multiple elastic 276 scattering which may affect the measured kinematic intensities (Fujiwara, 1959; Cowley, 1995). 277 For any typical hydrated protein crystal, these effects are suggested to be far less detrimental to data quality compared to inelastic scattering (Latychevskaia & Abrahams, 2019; Martynowycz, 278 Clabbers, Unge et al., 2021). Dynamical structure refinement can enhance the localization of 279 hydrogen atoms in small molecule structures (Palatinus et al., 2017), but its implementation is 280 computationally expensive and has yet to be extended to macromolecules that include bulk 281 282 solvent that cannot be modeled. In recent experiments, recording MicroED data using a direct electron detector in electron counting mode significantly improved data quality, and we expect 283 further benefits from faster readout and better electron-counting algorithms using electron-event 284 285 representation (Guo et al., 2020; Nakane et al., 2020).

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290 Crystallization and sample preparation

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Crystalline lamellae of triclinic lysozyme were prepared as described previously (Martynowycz, 292 Clabbers, Hattne et al., 2021). Briefly, crystals of hen egg-white lysozyme (Gallus gallus) were 293 grown by dissolving 10 mg/ml protein in a solution of 0.2 M sodium nitrate and 50 mM sodium 294 acetate at pH 4.5. After incubation overnight at 4 °C an opaque suspension was observed. After 295 further incubation for one week at room temperature a crystalline slurry appeared containing 296 297 microcrystals. Samples were prepared by depositing 3 µl of the crystalline slurry onto a glowdischarged EM grid (Quantifoil, Cu 200 mesh, R2/2 holey carbon). Excess liquid was blotted 298 away and the sample was vitrified using a Leica GP2 vitrification robot. Grids were transferred 299 to an Aquilos dual-beam FIB/SEM (Thermo Fisher) and crystals were milled to lamellae with an 300 301 optimal thickness of approximately 300 nm as described previously (Martynowycz, Clabbers, Hattne et al., 2021; Martynowycz, Clabbers, Unge et al., 2021). 302

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- 304 Data collection and processing
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Electron-counted MicroED data were collected on a Titan Krios 3Gi TEM (Thermo Fisher) 306 operated at 300 kV as described previously (Martynowycz, Clabbers, Hattne et al., 2021). 307 Briefly, the TEM was set up for low exposure data collection using a 50 µm C2 aperture, spot 308 size 11, and a beam diameter of 25 µm. A 100 µm SA aperture was used, corresponding to an 309 area of 2 µm diameter on the specimen. Crystal lamellae were continuously rotated over a range 310 311 of 84° at a rotation speed of 0.2°/s over 420s with a total exposure of approximately 0.64 e⁻.Å⁻² per dataset. Data were recorded on a Falcon 4 direct electron detector (Thermo Fisher) in 312 313 electron counting mode operating at an internal frame rate of 250 Hz. Data of 16 crystal lamellae

were integrated, scaled and merged using XDS (Kabsch, 2010) and AIMLESS (Evans & Murshudov, 2013). The structure was phased *ab initio* by placing a three-residue idealized α -helix fragment using PHASER (McCoy et al., 2007) followed by density modification in ACORN (Foadi et al., 2000). The entire structure was built automatically using BUCCANEER (Cowtan, 2006) and refined in *REFMAC5* (Murshudov et al., 2011) using electron scattering factors. Identification of hydrogen atoms The structure was manually inspected and remodeled using Coot (Emsley et al., 2010), and re-refined with REFMAC5 (Murshudov et al., 2011) using electron scattering factors. Hydrogen atoms were added in idealized riding positions. A hydrogen-only omit was calculated from the final structural model by REFMAC5 (Murshudov et al., 2011). Peaks in the $mF_0 - DF_c$ difference map at a threshold above 2.0σ were identified and listed using *PEAKMAX* in the CCP4 software package (Winn et al., 2011). Difference peaks that fell within 0.5 Å of the idealized distance for the known positions were assigned as hydrogen atoms. Figure preparation Figures were prepared using ChimeraX and assembled in powerpoint and photoshop. Data availability Coordinates and structure factors have been deposited to the PDB. Acknowledgements This study was supported by the National Institutes of Health P41GM136508. The Gonen laboratory is supported by funds from the Howard Hughes Medical Institute.

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478 Figure 1. Hydrogen atoms and bonding interactions in secondary structure elements. Difference 479 peaks for individual hydrogen atoms are displayed as green spheres with their σ values shown for (a) two 480 short anti-parallel β -strands (residues 42-45 and 51-54, respectively), and (b) an α -helix (residues 88-481 101). Hydrogen atoms were assigned from a hydrogen-only omit map for peaks at $\geq 2.0\sigma$ that are within 482 0.5 Å distance from their idealized position. Hydrogen bonding interactions are indicated by dashed black 483 lines and their respective bond distances and angles are listed in Table 2. Electrostatic potential 2mFo-484 DFc maps are contoured at 4.0 σ (blue) and mFo-DFc difference maps are shown at 2.5 σ (green and red 485 for positive and negative, respectively). Carbon atoms are shown in brown, nitrogen in blue, and oxygen in red. 486 487

488 Figure 2. Hydrogen atoms and hydrogen-bond networks. Hydrogen atoms (green difference peaks) 489 are shown with their σ values for the side chains of different residues and for several water molecules. 490 Hydrogen atoms were assigned from a hydrogen-only omit map for peaks at $\geq 2.0\sigma$ and within 0.5Å from 491 their idealized positions. (a) Strong difference peaks indicate hydrogen atom positions for His15, as well 492 as a possible hydrogen bond interaction between His15-NE2 and a neighboring Thr89-OG1. The histidine 493 residue appears to be protonated at ND1 which is consistent with pH 4.5 of the crystallization condition. 494 (b) Hydrogen atoms are indicated by difference peaks for two residues, as well as a potential hydrogen 495 bonding interaction between Tyr53-OH and Asp66-OD1. (c) Acidic side-chains showing well resolved atoms. Strong difference peaks for side chain hydrogen atoms can be observed in asparagine and 496 497 glutamine residues. (d) Illustration of a hydrogen bonding network involving water molecules and several 498 protein residues. The inset shows the hydrogen bond distances for the water molecules. Electrostatic 499 potential 2mFo-DFc maps are contoured at (a) 2.5σ (blue) and (b-d) at 3.0σ (blue), mFo-DFc difference 500 maps are shown at 2.3σ (green and red for positive and negative, respectively). Carbon atoms are shown 501 in brown, nitrogen in blue, and oxygen in red.

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Figure 3. Hydrogen bond distances for macromolecular MicroED data. Hydrogen bond distances in Å are shown as histogram plots with a normal distribution fitted to the data. Idealized hydrogen bond lengths between electron cloud centroids used in X-ray diffraction are indicated by a teal dotted line, idealized inter-nuclei hydrogen bond lengths used in neutron diffraction are indicated by the orange dotted line (see also Table 1, Supplementary Tables 2, 3, 6–8) (Gruene *et al.*, 2014).

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522 Table 1. Hydrogen atoms and mean observed hydrogen bond lengths

Hydrogen bonds	No. observations	X-H (Å) ^a	X-H _{X-ray} (Å) ^b	X-H _{Neutron} (Å) ^c
C _a .H	61	1.11(13)	0.98	1.10
C _{sidechain} -H	14	1.25(18)	0.98	1.10
C _{aromatic} -H	17	1.13(12)	0.93	1.08
C-H ₂	99	1.17(15)	0.97	1.09
C-H ₃	77	1.09(17)	0.96	1.06
N-H	44	1.02(16)	0.86	1.01
N-H•••O	38	1.05(15)	0.86	1.01
N-H ₂	13	1.08(21)	0.89	1.03
N-H ₃	3	1.13(11)	0.86	1.01
О-Н	10	1.13(18)	0.82	0.98

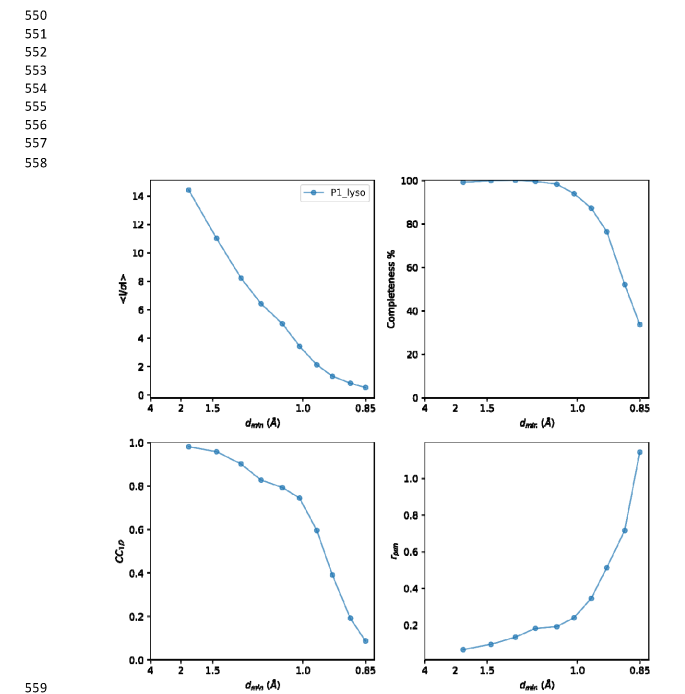
^a Mean observed hydrogen bond lengths measured for hydrogen atoms difference peaks at ≥2.0σ, standard deviations are listed in parenthesis. Values for individual hydrogen bond distances are listed in Supplementary Tables 2–10

^b Idealized hydrogen bond lengths between electron cloud centroids used in X-ray diffraction (Gruene *et al.*, 2014)

^c Idealized inter-nuclei hydrogen bond lengths used in neutron diffraction (Gruene *et al.*, 2014)

546 Table 2. Hydrogen bond distances and angles for secondary structure

Donor-H•••Acceptor	Diff. peak σ	D-H (Å)	H•••A (Å)	D•••A (Å)	D-H•••A (°)
β-strands					
Asn ⁴⁴ -N-H•••Asp ⁵² -O	3.13	1.41	1.46	2.86	170.74
Asp ⁵² -N-H•••Asn ⁴⁴ -O	4.33	0.90	1.93	2.75	149.88
Gly ⁵⁴ -N-H•••Thr ⁴³ -O	4.48	1.10	1.59	2.67	168.44
α-helices					
Ala ¹¹ -N-H•••Glu ⁷ -O	2.58	0.78	2.40	2.89	122.19
Met ¹² -N-H•••Leu ⁸ -O	2.44	0.90	1.87	2.70	153.21
Lys ¹³ -N-H•••Ala ⁹ -O	3.44	1.21	1.59	2.76	162.32
Arg ¹⁴ -N-H•••Ala ¹⁰ -O	2.39	1.08	2.02	2.82	128.88
Val ²⁹ -N-H•••Leu ²⁵ -O	2.09	0.77	2.39	2.94	129.35
Cys ³⁰ -N-H•••Gly ²⁶ -O	2.45	1.00	1.85	2.74	146.82
Ala ³¹ -N-H•••Asn ²⁷ -O	3.23	1.10	1.87	2.79	138.11
Lys ³³ -N-H•••Val ²⁹ -O	3.03	0.86	2.01	2.81	153.99
Phe ³⁴ -N-H•••Cys ³⁰ -O	3.21	0.79	2.18	2.95	169.63
Val ⁹² -N-H•••Ile ⁸⁸ -O	3.00	0.98	1.79	2.77	174.55
Asn ⁹³ -N-H•••Thr ⁸⁹ -O	3.67	0.96	1.79	2.74	168.88
Ala ⁹⁵ -N-H•••Ser ⁹¹ -O	2.74	0.87	1.86	2.73	172.72
Lys ⁹⁶ -N-H•••Val ⁹² -O	4.21	1.05	1.76	2.81	174.10
Ile ⁹⁸ -N-H•••Cys ⁹⁴ -O	3.04	1.13	1.72	3.15	152.39
Val ⁹⁹ -N-H•••Ala ⁹⁵ -O	3.06	1.09	1.95	3.03	171.83



Supplementary Figure 1. Merging statistics for triclinic lysozyme. Crystallographic quality indicators and data completeness plotted as function of resolution for triclinic lysozyme at 0.87 Å resolution (see Supplementary Table 1).

577 Supplementary Table 1. MicroED Data collection and refinement statistics

Data collection [*]	
Wavelength	0.0197
No. of crystals	16
Space group	P1
Cell dimensions	
<i>a</i> , <i>b</i> , <i>c</i> (Å)	26.42, 30.72, 33.01
<i>α</i> , <i>β</i> , <i>γ</i> (°)	88.32, 109.10, 112.08
Resolution (Å)	16.05-0.87 (0.90-0.87)**
Observed reflections	569407 (5797)
Unique reflections	64986 (2783)
Multiplicity	8.8 (2.1)
Completeness (%)	87.55 (37.64)
R _{merge}	0.236 (1.035)
R _{meas}	0.248 (1.409)
R _{pim}	0.073 (0.945)
Mean I/ $\sigma(I)$	6.23 (0.66)
CC _{1/2}	0.990 (0.147)
Refinement	
No. of reflections	64974
No. of reflections used for R_{free}	3168
R_{work} / R_{free}	0.197 / 0.220
No. of atoms	1216
Proteins	1081
Ligand	16
Water	119
R.m.s. deviations	
Bond lengths (Å)	0.034
Bond angles (°)	2.447
Mean <i>B</i> -factor ($Å^2$)	11.98
Ramachandran	

Favored (%)	96.19
Allowed (%)	3.81
Outliers (%)	0.00
Rotamer outliers (%)	2.33

^{*}Data from Martynowycz, Clabbers, Hattne *et al.*, 2021.

^{**}Values in parentheses are for highest-resolution shell.

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583 Supplementary Table 2. Bond distances and angles for hydrogen bonding interactions

Donor-H•••Acceptor	Diff. peak σ	D-H (Å)	H•••A (Å)	D•••A (Å)	D-H•••A (°)
Phe ³ -N-H•••Phe ³⁸ -O	2.50	1.18	1.67	2.81	160.72
Ala ¹¹ -N-H•••Glu ⁷ -O	2.58	0.78	2.40	2.89	122.19
Met ¹² -N-H•••Leu ⁸ -O	2.44	0.90	1.87	2.70	153.21
Lys ¹³ -N-H•••Ala ⁹ -O	3.44	1.21	1.59	2.76	162.32
Arg ¹⁴ -N-H•••Ala ¹⁰ -O	2.39	1.08	2.02	2.82	128.88
His ¹⁵ -NE2-H•••Thr ⁸⁹ -OG1	2.39	1.27	1.33	2.59	162.48
Leu ¹⁷ -N-H•••Met ¹² -O	2.91	1.11	1.92	2.99	162.18
Tyr ²⁰ -N-H•••Leu ¹⁷ -O	2.78	1.06	1.98	2.92	146.41
Val ²⁹ -N-H•••Leu ²⁵ -O	2.09	0.77	2.39	2.94	129.35
Cys ³⁰ -N-H•••Gly ²⁶ -O	2.45	1.00	1.85	2.74	146.82
Ala ³¹ -N-H•••Asn ²⁷ -O	3.23	1.10	1.87	2.79	138.11
Lys ³³ -N-H•••Val ²⁹ -O	3.03	0.86	2.01	2.81	153.99
Phe ³⁴ -N-H•••Cys ³⁰ -O	3.21	0.79	2.18	2.95	169.63
Thr ⁴⁰ -N-H•••Lys ¹ -O	3.03	1.08	1.77	2.75	147.63
Ala ⁴² -N-H•••Asn ³⁹ -O	3.27	1.20	1.77	2.89	154.32
Asn ⁴⁴ -N-H•••Asp ⁵² -O	3.12	1.41	1.46	2.86	170.74
Asn ⁴⁶ -N-H•••Ser ⁵⁰ -O	2.56	1.21	1.63	2.73	148.76
Asp ⁵² -N-H•••Asn ⁴⁴ -O	4.33	0.90	1.93	2.75	149.88
Tyr ⁵³ -O-H•••Asp ⁶⁶ -OD2	3.44	1.39	1.25	2.59	157.86
Gly ⁵⁴ -N-H•••Thr ⁴³ -O	4.48	1.10	1.59	2.67	168.44
Gln ⁵⁷ -N-H•••Gly ⁵⁴ -O	3.01	1.11	1.83	2.84	150.17
Asn ⁶⁵ -N-H•••Leu ⁷⁸ -O	2.74	0.91	1.87	2.77	159.36
Gly ⁶⁷ -N-H•••Asn ⁶⁵ -O	3.90	1.13	1.85	2.90	152.42
Arg ⁷³ -N-H•••Arg ⁶¹ -O	3.77	1.17	1.74	2.80	147.11
Leu ⁷⁵ -N-H•••Trp ⁶² -O	2.86	0.84	1.93	2.75	161.86
Ser ⁸¹ -N-H•••NO3 ²⁰¹ -O	2.92	1.07	1.8	2.76	146.94
Ala ⁸² -N-H•••Pro ⁷⁰ -O	2.16	1.18	1.71	2.79	148.60
Leu ⁸³ -N-H•••Cys ⁸⁰ -O	2.32	1.04	1.77	2.73	150.33
Leu ⁸⁴ -N-H•••Ser ⁸¹ -O	2.99	1.03	1.92	2.93	164.49
Val ⁹² -N-H•••Ile ⁸⁸ -O	3.00	0.98	1.79	2.77	174.55
Asn ⁹³ -N-H•••Thr ⁸⁹ -O	3.67	0.96	1.79	2.74	168.89
Ala ⁹⁵ -N-H•••Ser ⁹¹ -O	2.74	0.87	1.86	2.73	172.72

Lys ⁹⁶ -N-H•••Val ⁹² -O	4.21	1.05	1.76	2.81	174.10
Ile ⁹⁸ -N-H•••Cys ⁹⁴ -O	3.04	1.13	1.72	3.15	152.39
Val ⁹⁹ -N-H•••Ala ⁹⁵ -O	3.06	1.09	1.95	3.03	171.83
Trp ¹⁰⁸ -N-H•••Met ¹⁰⁵ -O	3.10	1.26	1.64	2.82	152.36
Trp ¹¹¹ -NE1-H•••Asn ²⁷ -OD1	3.39	1.02	1.79	2.71	169.48
Arg ¹¹⁴ -N-H•••Arg ¹¹⁰ -O	2.27	1.11	1.98	2.76	124.52
Trp ¹²³ -N-H•••Val ¹²⁰ -O	2.45	1.04	1.83	2.84	161.33

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586 Supplementary Table 3. Hydrogen bond distances for Ca-H

Residue	Name	Atom	Diff. peak σ	X-H (Å)
2	Val	CA	3.42	1.15
3	Phe	CA	3.46	1.26
4	Gly	CA	3.18	1.14
4	Gly	CA	2.39	0.98
5	Arg	CA	2.59	1.27
7	Glu	CA	4.16	1.06
9	Ala	CA	3.79	1.08
10	Ala	CA	2.74	1.00
11	Ala	CA	2.78	0.81
12	Met	CA	2.74	1.11
13	Lys	CA	2.01	1.20
16	Gly	CA	2.60	0.89
18	Asp	CA	2.26	1.45
19	Asn	CA	2.81	1.43
20	Tyr	CA	3.17	1.10
21	Arg	CA	3.12	1.29
22	Gly	CA	2.77	1.18
24	Ser	CA	3.96	1.10
25	Leu	CA	3.06	1.19
27	Asn	CA	3.22	1.16
28	Trp	CA	3.34	1.10
29	Val	CA	2.42	1.10
32	Ala	CA	2.39	1.05
33	Lys	CA	2.71	1.05
38	Phe	CA	2.41	1.18
39	Asn	CA	2.96	0.99
42	Ala	CA	2.17	0.83
44	Asn	CA	2.16	0.90
45	Arg	CA	2.81	0.98
51	Thr	CA	3.55	0.90
52	Asp	CA	3.04	1.20
53	Tyr	CA	2.27	1.14
54	Gly	CA	3.43	1.17
55	Ile	CA	2.63	1.12
56	Ile	CA	2.85	1.25
57	Asn	CA	2.89	1.23
58	Ile	CA	3.49	0.97
59	Asn	CA	3.51	1.06
62	Trp	CA	2.78	1.18
63	Trp	CA	2.29	1.25

69	Thr	CA	2.61	1.00
70	Pro	CA	2.91	1.06
74	Asn	CA	3.22	1.03
75	Leu	CA	2.65	1.13
76	Cys	CA	2.76	1.08
77	Asn	CA	2.87	1.13
82	Ala	CA	3.27	1.09
88	Ile	CA	2.64	1.41
91	Ser	CA	4.18	1.40
94	Cys	CA	2.89	1.02
96	Lys	CA	2.80	1.05
97	Lys	CA	2.38	1.24
99	Val	CA	3.06	1.15
103	Asn	CA	2.14	1.17
108	Trp	CA	4.03	0.94
110	Ala	CA	3.41	1.06
111	Trp	CA	2.44	1.10
116	Lys	CA	2.93	1.11
117	Gly	CA	4.02	1.11
119	Asp	CA	3.19	1.07
123	Trp	CA	2.60	1.14

622 Supplementary Table 4. Hydrogen bond distances for side chain C-H

Residue	Name	Atom	Diff. peak σ	X-H (Å)
8	Leu	CG	2.50	0.94
15	His	CE1	2.50	1.45
40	Thr	CB	2.64	1.51
56	Leu	CG	3.74	1.22
58	Ile	CB	3.35	1.18
63	Trp	CD1	3.06	1.39
69	Thr	CB	3.08	0.99
84	Leu	CG	2.88	1.29
89	Thr	CB	3.24	1.25
98	Ile	CB	2.19	1.06
108	Trp	CD1	2.84	1.25
109	Val	CB	2.36	1.51
118	Thr	CB	3.39	1.29
123	Trp	CD1	2.44	1.19

653 Supplementary Table 5. Hydrogen bond distances for aromatic C-H

Residue	Name	Atom	Diff. peak σ	X-H (Å)
3	Phe	CE2	3.12	1.05
20	Tyr	CE1	3.09	0.99
23	Tyr	CD2	4.18	1.01
23	Tyr	CD1	2.81	1.11
23	Tyr	CE1	2.15	1.32
38	Phe	CZ	3.97	1.13
38	Phe	CE2	2.56	1.32
38	Phe	CD1	2.11	1.03
53	Tyr	CD1	3.52	1.18
53	Tyr	CE2	3.39	1.16
63	Trp	CZ3	3.51	1.00
63	Trp	CE3	3.19	1.22
108	Trp	CE3	3.86	1.07
108	Trp	CH2	3.47	1.01
108	Trp	CZ3	2.42	1.31
111	Trp	CH2	3.52	1.24
111	Trp	CZ2	2.30	1.02

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Supplementary Table 6. Hydrogen bond distances for CH2 683

Diff. peak σ X-H (Å) Residue Name Atom CE 3.20 1.41 1 Lys 1 Lys CE 2.30 1.17 1 Lys CG 3.15 1.05 CB 3.12 1 1.16 Lys 3 Phe CB 3.37 0.98 3 2.43 1.14 Phe CB 5 Arg CB 3.12 1.23 5 CB 2.81 1.46 Arg 5 CG 2.20 1.14 Arg 2.37 6 CB 1.04 Cys 7 Glu CB 2.42 1.58 8 Leu CB 2.45 1.03 8 Leu CB 2.43 1.10 12 CG 3.12 1.18 Met CG 1.00 14 3.82 Arg 15 His CB 2.07 1.16 18 Asp CB 2.84 1.00 18 Asp CB 2.29 1.21 19 CB 3.04 1.01 Asn 19 CB 2.48 1.06 Asn 2.43 20 Tyr CB 1.06 2.20 20 Tyr CB 1.07 CG 3.36 1.17 21 Arg 21 CB 2.38 1.16 Arg 21 CB 2.16 1.47 Arg 23 CB 2.80 1.14 Trp 25 Leu CB 3.22 1.08 27 CB 2.69 0.91 Asn 28 Trp CB 2.19 1.06 3.45 1.07 33 CE Lys 33 CG 3.11 1.08 Lys 33 CG 2.27 Lys 1.13 34 Phe CB 4.37 1.18 Glu CG 3.11 1.17 35 35 Glu CG 2.53 1.34 CB 3.51 1.09 36 Ser CB 1.51 36 Ser 2.63 38 Phe CB 3.14 1.06 CB 41 Gln 3.12 1.19 41 Gln CB 2.76 1.00

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41	Gln	CG	2.48	1.10
48	Asp	CB	2.11	1.29
57	Gln	CB	3.45	1.18
57	Gln	СВ	2.49	1.19
58	Ile	CG1	3.79	1.23
58	Ile	CG1	2.79	1.12
59	Asn	CB	3.11	1.15
59	Asn	CB	2.52	1.06
60	Ser	CB	3.67	1.30
61		CB	4.18	1.04
61	Arg	CB	2.59	0.91
61	Arg	CG	2.39	1.47
62	Arg	CB	3.11	1.25
63	Trp Trp	CB	2.97	1.29
63	Trp	CB	2.37	1.29
64	Cys	CB	2.54	1.29
66	Asp	CB	2.55	1.19
66	Asp	CB	2.09	1.19
68	-	CB	4.31	1.16
70	Arg Pro	CB	2.76	1.10
70	Ser	CB	2.04	1.14
72	Arg	CB	2.22	1.14
73	Asn	CB	2.22	1.16
74	Asn	CB	2.40	1.10
75	Leu	CB	2.82	1.35
76	Cys	CB	3.12	0.76
76	Cys	CB	3.07	1.21
77	Asn	CB	2.76	1.08
78	Ile	CG1	2.13	1.05
80	Cys	CB	3.14	0.99
80	Cys	CB	2.60	1.31
83	Leu	CB	2.57	1.01
84	Leu	CB	2.72	1.23
85	Ser	CB	2.37	1.28
86	Ser	CB	2.91	1.22
87	Asp	СВ	3.81	1.20
88	Ile	CG1	2.16	1.34
91	Ser	СВ	2.86	0.95
91	Ser	СВ	2.10	1.16
93	Asn	СВ	3.61	1.40
93	Asn	СВ	2.59	1.15
94	Cys	СВ	3.46	1.08
94	Cys	CB	2.50	1.20
96	Lys	CB	3.05	0.90
97	Lys	CD	4.05	1.25
97	Lys	CG	3.62	1.07
97	Lys	CG	2.88	1.26
97	Lys	CD2	2.97	1.30
97	Lys	CD2	2.65	1.18
97	Lys	CB	2.60	0.82
100	Ser	CB	3.19	1.24
105	Met	CB	2.62	1.42
105	Met	CG	2.35	1.23

108	Trp	CB	3.41	1.17
114	Arg	CB	2.26	1.16
116	Lys	CE	3.61	1.23
121	Gln	CB	2.77	1.04
121	Gln	СВ	2.50	1.23
123	Trp	CB	2.11	1.41

688 Supplementary Table 7. Hydrogen bond distances for CH₃ 689

Residue	Name	Atom	Diff. peak σ	X-H (Å)
2	Val	CG1	3.42	0.94
2	Val	CG1	2.23	1.15
2	Val	CG2	3.04	1.31
2	Val	CG2	2.63	1.11
8	Leu	CD1	3.59	1.20
8	Leu	CD1	3.12	0.99
9	Ala	СВ	2.93	1.00
10	Ala	СВ	3.73	1.12
10	Ala	СВ	2.89	1.54
10	Ala	СВ	2.56	1.03
12	Met	CE	3.02	1.14
12	Met	CE	2.27	1.11
17	Leu	CD2	2.75	0.99
17	Leu	CD1	2.52	0.90
25	Leu	CD2	3.61	0.87
25	Leu	CD2	3.04	1.12
25	Leu	CD1	2.31	1.03
29	Val	CG1	3.91	0.99
29	Val	CG1	2.92	1.40
29	Val	CG2	2.42	1.06
31	Ala	CB	2.64	0.91
31	Ala	СВ	2.47	1.05
31	Ala	CB	2.53	0.87
32	Ala	CB	2.63	0.90
40	Thr	CG2	2.99	1.28
42	Ala	CB	2.71	0.94
51	Thr	CG2	3.29	1.18
51	Thr	CG2	2.31	1.12
55	Ile	CD1	2.71	1.12
55	Ile	CD1	2.30	1.20
55	Ile	CG2	2.54	0.97
56	Leu	CD1	3.20	1.35
56	Leu	CD1	2.52	1.06
56	Leu	CD1	2.56	1.31
56	Leu	CD2	2.67	0.99
56	Leu	CD2	2.47	1.14
58	Ile	CD1	2.82	1.09
58	Ile	CD1	2.54	1.09
58	Ile	CG2	3.20	1.26

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58	Ile	CG2	3.00	0.87
69	Thr	CG2	2.05	1.21
75	Leu	CD1	3.45	1.08
75	Leu	CD1	2.35	0.89
75	Leu	CD2	2.27	0.95
78	Ile	CG2	3.55	1.27
78	Ile	CG2	2.34	1.25
82	Ala	CB	3.39	0.89
82	Ala	CB	3.12	1.04
83	Leu	CD2	2.34	1.36
84	Leu	CD2	3.58	1.18
84	Leu	CD1	2.80	1.02
88	Ile	CD1	3.97	1.27
88	Ile	CD1	2.12	1.36
88	Ile	CG2	2.43	1.09
88	Ile	CG2	2.21	1.16
90	Ala	СВ	4.32	1.13
92	Val	CG2	3.67	0.92
92	Val	CG2	3.34	0.81
92	Val	CG2	3.08	1.02
95	Ala	CB	2.63	1.02
95	Ala	CB	2.19	0.96
98	Ile	CD1	3.03	1.19
98	Ile	CD1	2.15	0.79
99	Val	CG2	2.93	1.13
99	Val	CG2	2.79	1.07
105	Met	CE	3.46	1.16
107	Ala	CB	2.74	1.32
110	Ala	CB	2.43	1.39
110	Ala	CB	2.03	0.96
118	Thr	CG2	2.09	1.03
120	Val	CG2	3.01	1.08
120	Val	CG2	2.09	0.76
120	Val	CG1	2.42	1.59
122	Ala	CB	2.64	1.17
122	Ala	CB	2.03	1.14
124	Ile	CG2	2.79	0.98
124	Ile	CG2	2.67	0.65

Supplementary Table 8. Hydrogen bond distances for N-H 712

Residue	Name	Atom	Diff. peak σ	X-H (Å)
4	Gly	N	3.10	0.94
6	Cys	N	2.34	1.24
7	Glu	N	2.72	0.87
14	Arg	NE1	2.47	1.19
18	Asp	N	2.90	0.91
21	Arg	N	3.12	1.26
22	Gly	N	3.94	0.80
23	Tyr	N	4.43	1.13
25	Leu	N	2.62	0.93
26	Gly	N	3.30	1.00
37	Asn	N	2.66	1.09
38	Phe	N	2.98	1.18
39	Asn	N	3.20	1.14
45	Arg	N	2.35	0.75
49	Gly	N	2.70	0.98
55	Ile	N	2.70	0.88
56	Ile	N	2.72	1.21
58	Ile	N	3.27	1.08
60	Ser	N	3.37	0.91
61	Arg	N	4.43	0.94
62	Trp	N	2.96	0.81
62	Trp	NE1	2.35	1.11
71	Gly	N	2.21	1.12
72	Ser	N	2.27	1.07
74	Asn	N	2.06	0.73
76	Cys	N	2.76	1.19
78	Ile	N	2.58	0.85
80	Cys	N	2.11	0.98
86	Ser	N	2.12	1.12
88	Ile	N	2.09	0.91
89	Thr	N	4.00	1.14
90	Ala	N	3.27	1.13
91	Ser	N	2.14	0.99
104	Gly	N	2.87	1.24
105	Met	N	2.64	1.25
107	Ala	N	3.15	0.71
108	Trp	NE1	2.37	1.14
109	Val	N	3.68	0.81
111	Trp	Ν	3.22	1.21

118	Thr	N	4.54	1.02
119	Asp	N	2.50	1.26
120	Val	N	3.86	1.16
121	Asn	N	2.50	0.88
124	Ile	Ν	3.53	0.84

Supplementary Table 9. Hydrogen bond distances for N-H₂ 719

Residue	Name	Atom	Diff. peak σ	X-H (Å)
5	Arg	NH1	2.43	1.14
14	Arg	NH2	2.96	0.76
14	Arg	NH1	2.93	1.26
19	Asn	ND2	2.63	1.31
19	Asn	ND2	2.37	1.14
27	Asn	ND2	3.07	0.87
27	Asn	ND2	2.00	1.13
39	Asn	ND2	2.18	1.17
57	Gln	NE2	2.80	1.00
59	Asn	ND2	2.21	0.75
62	Arg	NH1	2.42	1.10
74	Asn	ND2	3.40	1.50
114	Arg	NH1	2.07	0.92

722 Supplementary Table 10. Hydrogen bond distances for N-H₃

Residue	Name	Atom	Diff. peak σ	X-H (Å)
1	Lys	NZ	3.32	1.24
33	Lys	NZ	2.70	1.13
97	Lys	NZ	2.96	1.02

