# Structure of human DPPA3 bound to the UHRF1 PHD finger reveals its functional and structural differences from mouse DPPA3

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# 19 Abstract

DNA methylation maintenance is essential for cell fate inheritance. In differentiated cells, this 20 involves orchestrated actions of DNMT1 and UHRF1. In mice, the high-affinity binding of 21 DPPA3 to the UHRF1 PHD finger regulates UHRF1 chromatin dissociation and cytosolic 22 localization, which is required for oocyte maturation and early embryo development. However, 23 24 the human DPPA3 ortholog functions during these stages remain unclear. Here, we report the 25 structural basis for human DPPA3 binding to the UHRF1 PHD finger. The conserved human DPPA3 <sup>85</sup>VRT<sup>87</sup> motif binds to the acidic surface of UHRF1 PHD finger, whereas mouse 26 DPPA3 binding additionally utilizes two unique  $\alpha$ -helices. The binding affinity of human 27 DPPA3 for the UHRF1 PHD finger was weaker than that of mouse DPPA3. Consequently, 28 human DPPA3, unlike mouse DPPA3, failed to inhibit UHRF1 chromatin binding and DNA 29 remethylation in Xenopus egg extracts effectively. Our data provide novel insights into the 30 distinct function and structure of human DPPA3. 31

# 33 Introduction

DNA methylation, a cytosine methylation at the 5th carbon atom in a CpG sequence, is a major 34 epigenetic mark that regulates diverse biological processes, including cell-type-specific gene 35 expression, retrotransposon silencing, X-chromosome inactivation, genome imprinting, and 36 carcinogenesis<sup>1,2</sup>. Once DNA methylation patterns are established during cell differentiation, 37 they are faithfully inherited after each replication, to maintain cell identity<sup>3,4</sup>. DNMT1, a 38 maintenance DNA methyltransferase, and UHRF1 (ubiquitin-like PHD and RING finger 39 domain-containing protein 1, also known as Np95/ICBP90), a ubiquitin E3-ligase and recruiter 40 of DNMT1, play pivotal roles in maintaining DNA methylation<sup>5-8</sup>. During this process, the 41 UHRF1 SET- and RING-associated (SRA) domain specifically binds to hemi-methylated 42 DNA<sup>9-11</sup>, and UHRF1 ubiquitinates histone H3 or PAF15 (PCNA-associated factor 15) using 43 a plant homeodomain (PHD) finger for recognition, and ubiquitin-like (UBL) and really 44 interesting new gene (RING) domains for multiple mono-ubiquitination<sup>12-17</sup>. Ubiquitinated 45 histone H3 and PAF15 recruit DNMT1 to the late- and early replicating domains, 46 respectively<sup>17–19</sup>, and stimulate the methyltransferase activity of DNMT1<sup>14,20</sup>. 47

In addition to its well-established role in DNA methylation maintenance, UHRF1 has emerged 48 as a factor in oocyte and preimplantation embryo development<sup>21-23</sup>. A maternal factor, 49 developmental pluripotency-associated 3 (DPPA3), also known as Stella/PGC7, has been 50 identified in mice as a strict inhibitor of chromatin binding of UHRF1 and regulation of its 51 cytosolic localization, in cooperation with exportin-1<sup>24-26</sup>. Expression of mouse DPPA3 52 (mDPPA3), an intrinsically disordered protein, is restricted to primordial germ cells, oocytes, 53 and preimplantation embryos<sup>24,27,28</sup>. mDPPA3 plays an important role in the formation of 54 oocyte-specific DNA methylation patterns by preventing excessive *de novo* DNA methylation 55 mediated by UHRF1<sup>24</sup>. Using nuclear magnetic resonance (NMR) solution structural analysis 56 of mouse the UHRF1 PHD finger (mPHD) bound to mDPPA3, we recently revealed that the 57 C-terminal region of mDPPA3 binds to mPHD utilizing a VRT motif at residues 88-90 58

 $^{(88}$ VRT<sup>90</sup>), which is conserved in the motifs of other binding partners, histone H3 <sup>1</sup>ART<sup>3</sup> and PAF15 <sup>1</sup>VRT<sup>3</sup> with two subsequent α-helices unique to mDPPA3<sup>29</sup>. Owing to this multifaceted interaction, the binding affinity of mDPPA3 to mPHD ( $K_D$  of 0.0277 µM) is significant stronger than those of histone H3 and PAF15 ( $K_D$  of 1.59 µM and 3.52 µM, respectively), indicating that the mechanism by which mDPPA3 inhibits chromatin-binding of UHRF1 involves the competitive binding of between mDPPA3 and histone H3/PAF15 to UHRF1<sup>29</sup>.

The biological functions of mDPPA3 as a demethylation factor and UHRF1-inihibitor in oocyte 65 and preimplantation embryos have been extensively studied in mouse models. A recent report 66 has shown that UHRF1 is enriched in the cytoplasmic lattices of human oocytes<sup>30</sup>. However, it 67 is unclear if the biological function of mDPPA3 is conserved in human DPPA3 (hDPPA3), and 68 its role in human oocytes and preimplantation embryos is unknown. Two  $\alpha$ -helices in mDPPA3 69 70 which are induced upon binding to mPHD has been shown to be required for the interaction 71 with mUHRF1<sup>29</sup>. However, the amino acid sequences corresponding to these helices are poorly conserved between human and mouse DPPA3 (Figure 1a), which raises a question of whether 72 73 hDPPA3 also binds to the hUHRF1 PHD finger in a manner similar to their mouse counterparts, and whether hDPPA3 can inhibit chromatin binding of UHRF1. 74

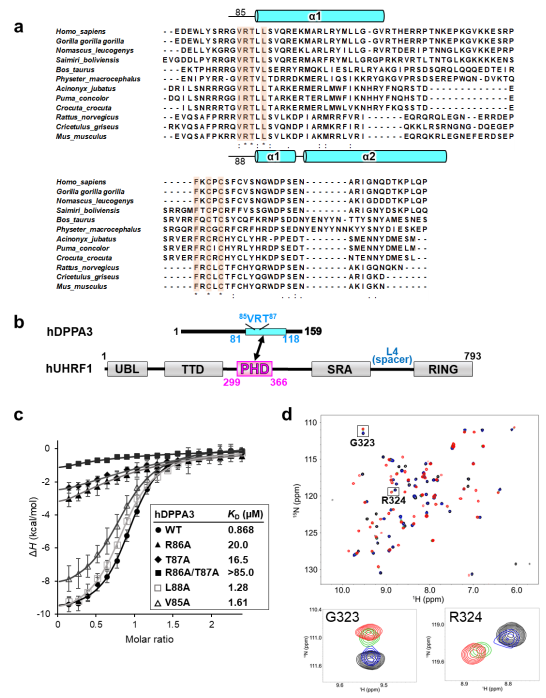
In this study, we determined the crystal structure of the human UHRF1 PHD finger complexed 75 with the C-terminal hDPPA3 fragment. The structure clearly showed that the binding mode of 76 hDPPA3 to the human UHRF1 PHD finger differs markedly from that of the mouse proteins 77 and explains why hDPPA3 binds to the human UHRF1 PHD finger with low binding affinity, 78 comparable to the binding of histone H3 and PAF15. Biochemical assays using *Xenopus* egg 79 80 extracts demonstrated that the inhibitory effect of hDPPA3 on chromatin-binding of UHRF1 is 81 relatively modest compared to the strong inhibition by mouse DPPA3. Our findings shed light on the unexpected role of hDPPA3 in epigenetic regulation during early embryonic 82 development, which differs from the evidence in mice. 83

# 84 **RESULTS**

## 85 Interaction between hDPPA3 and hUHRF1 PHD finger

Our previous NMR structural analysis of mDPPA3 complexed with mUHRF1 PHD (mPHD) 86 revealed that residues 85–118 of mDPPA3 are essential for its interaction with mPHD (Figure 87 1a and 2b)<sup>29</sup>. Thus, we identified the corresponding region of hDPPA3 by sequence alignment 88 (residues 81-118 [hDPPA3<sub>81-118</sub>] (Figure 1a and 1b), and evaluated whether this region binds 89 90 to the human UHRF1 PHD finger, residues 299–366 (hPHD). Isothermal titration calorimetry (ITC) demonstrated that hDPPA3<sub>81-118</sub> could bind to hPHD with a  $K_d$  of 0.868  $\mu$ M (Figure 1c), 91 which is approximately 30-fold weaker than the binding affinity between mDPPA3 and mPHD 92  $(K_{\rm d} = 0.0277 \ \mu {\rm M})^{29}$ . The binding affinity of hDPPA3 to hPHD<sub>81-118</sub> is comparable with the 93 previously reported binding affinity between hPHD and the histone H3 N-terminal tail 94 (residues 1–15;  $K_D = 1.7 \ \mu\text{M}$ ) or PAF15 (residues 1–10;  $K_D = 2.2 \ \mu\text{M}$ )<sup>17,31</sup>. To further 95 investigate the interactions at an atomic resolution, we performed NMR titration experiments. 96 We successfully assigned <sup>1</sup>H-<sup>15</sup>N heteronuclear single quantum coherence (HSQC) spectra to 97 <sup>15</sup>N]-hPHD in the free and complex states with non-labeled hDPPA3<sub>81-118</sub> (Supplementary 98 Figure 1a). The <sup>1</sup>H-<sup>15</sup>N HSQC spectra of [<sup>15</sup>N]-hPHD titrated with non-labeled hDPPA3<sub>81-118</sub> 99 showed that the HSQC signals shifted in the intermediate exchange regime on a chemical shift 100 timescale, supporting the modestly weak interaction between hDPPA3<sub>81-118</sub> and hPHD (Figure 101 1d). These data indicate that the binding of hDPPA3 to hPHD was not significantly stronger 102 than that of the other binding partners, histone H3 and PAF15. Chemical shift differences 103 (CSD) between the free and complexed states showed relatively large values for Asp330, 104 Met332, Asp337, Glu355, and Asp356, suggesting the contribution of the main chain of these 105 amino acid residues to the hPHD-hDPPA3 interaction (Supplementary Figure 1b). 106

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109 Figure 1: Characterization of the interaction between mUHRF1 and mDPPA3.

(a) Schematic of the domain composition of human UHRF1 and DPPA3. (b) Amino acid
 sequence alignment of C-terminal part of DPPA3. Secondary structures of mouse and human

112 DPPA3 are indicated based on PDB:7XGA and analysis of this study, respectively. (c) 113 Isothermal titration calorimetry measurements for hPHD and wild-type (WT)/mutants of

hDPPA3<sub>81-118</sub>. Superimposition of enthalpy change plots with standard deviations. Data were

presented as mean values for n = 3. (d) Overlay of <sup>1</sup>H-<sup>15</sup>N heteronuclear single quantum

116 coherence (HSQC) spectra of 30 µM mPHD showing chemical shift changes upon titration

with hDPPA3<sub>81-118</sub> of 0  $\mu$ M (black), 15  $\mu$ M (blue), 30  $\mu$ M (green), and 60  $\mu$ M (red). Square

regions inside the HSQC spectra were expanded (lower panels).

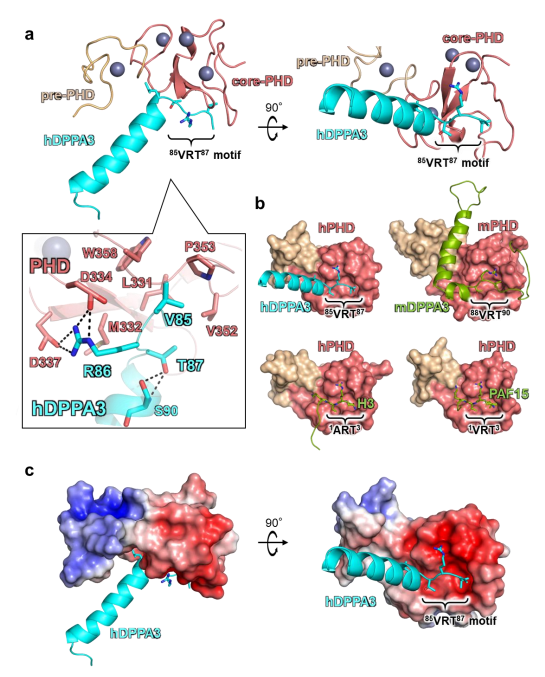
# 119 Crystal structure of hDPPA3 bound to hPHD

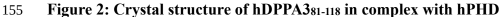
To reveal the molecular basis for the binding mode of hDPPA3 to hPHD, we determined the 120 crystal structure of hDPPA3<sub>81-118</sub> in complex with hPHD at a 2.4 Å resolution (Table 1). The 121 asymmetric unit contained one hPHD:hDPPA3<sub>81-118</sub> complex, and a  $2|F_0|-|F_c|$  map 122 corresponding to residues 299-363 of hPHD and residues 84-107 of hDPPA3<sub>81-118</sub> was 123 unambiguously observed (Supplementary Figure 2a). hPHD consists of pre- and core-PHD 124 domains that include three zinc finger motifs (Figure 2a)<sup>31</sup>. The structure of the hPHD moiety 125 in the complex with hDPPA3<sub>81-118</sub> was well-superimposed on apo-hPHD (PDB:3SOX, root 126 mean square deviation [RMSD] of Cα atoms with 0.848 Å) and those in the complex with 127 histone H3 (PDB:3ASL, RMSD: 0.795 Å) and PAF15 (PDB:6IIW, RMSD: 0.397 Å), implying 128 that the binding of hDPPA3 does not undergo conformational changes in hPHD 129 (Supplementary Figure 2b). 130

In contrast to the hPHD moiety, the binding mode of hDPPA3 shares both similarities and 131 dissimilarities with that of mDPPA3 (Figure 2a, 2b). The conserved VRT motif at residues 85-132 87 of hDPPA3<sub>81-118</sub> is accommodated on the acidic surface of hPHD, the binding site for <sup>1</sup>ART<sup>3</sup> 133 of histone H3, and <sup>1</sup>VRT<sup>3</sup> of PAF15, in a manner concordant with the motif in mDPPA3 134 (<sup>88</sup>VRT<sup>90</sup>) (Figure 2b). The side chain of Val85–hDPPA3<sub>81-118</sub> forms a hydrophobic interaction 135 with Leu331, Val352, Pro353, and Trp358 in hPHD (Figure 2a). The positively charged 136 guanidino group at Arg86 of hDPPA3<sub>81-118</sub> forms hydrogen bonds with the side chains of 137 Asp334 and Asp337 of hPHD (Figure 2a and 2c). The side chain methyl and hydroxyl groups 138 of Thr87 in hDPPA3<sub>81-118</sub> forms hydrophobic interactions with Leu331 and Val352 of hPHD 139 and hydrogen bonds with the main chain amide of Ser90 of hDPPA3 (Figure 2a). The latter 140 potentially functions as a helical cap for the N-terminus of the following  $\alpha$ -helix (Figure 2a). 141 Leu88 of hDPPA3<sub>81-118</sub> is surrounded by the side chains of Ala317, Gln330, Met332, and 142 Ala339 in hPHD, in which the side chain of Met332 functions as a separation between the side 143 chains of Arg86 and Leu88 of hDPPA3<sub>81-118</sub> (Figure 2a). 144

When mPHD binds to mDPPA3, the two  $\alpha$ -helices following the VRT motif of mDPPA3 form 145 an L-like shape, in which the long  $\alpha$ -helix binds to the shallow groove between the pre- and 146 core-PHD fingers (Figure 2b). However, the C-terminus of the <sup>85</sup>VRT<sup>87</sup> motif of hDPPA3<sub>81-118</sub> 147 forms a unique conformation that differed from that of mDPPA3. Residues 88–101 of hDPPA3 148 forms a four-turn single  $\alpha$ -helix, which is not kinked and markedly differs from mDPPA3 149 complexed with mPHD (Figure 2a and 2b). The contact area between the hPHD and hDPPA3 150 (ca. 449 Å<sup>2</sup>) was smaller than that of the mouse protein (ca. 1360 Å<sup>2</sup>)<sup>32</sup>, which is concordant 151 with the weaker dissociation constant of the human proteins. 152

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(a) Overall structure of hPHD:hDPPA3<sub>81-118</sub> complex. Pre-PHD, core-PHD and hDPPA3 are 156 depicted as gold, salmon, and cyan cartoon models, respectively. The conserved VRT motif in 157 hDPPA3 is displayed as a stick model. Inset shows the interaction between the VRT motif of 158 hDPPA3 and hPHD. The black dotted line represents a hydrogen bond. (b) Structural 159 comparison of hPHD:hDPPA3 (this study, upper left), mPHD:mDPPA3 (PDB: 7XGA, upper 160 right), hPHD:H3 (PDB: 3ASL, bottom left) and hPHD:PAF15 (PDB: 6IIW, bottom right) 161 complexes. mDPPA3, H3 and PAF15 are shown as a green cartoon model and VRT (ART) 162 motif are represented as stick model. (c) Electrostatic surface potential of hPHD calculated 163 with program APBS<sup>33</sup>. The red and blue surface colors represent negative and positive 164 charges, respectively. hPPA3 is depicted as a cyan cartoon. 165

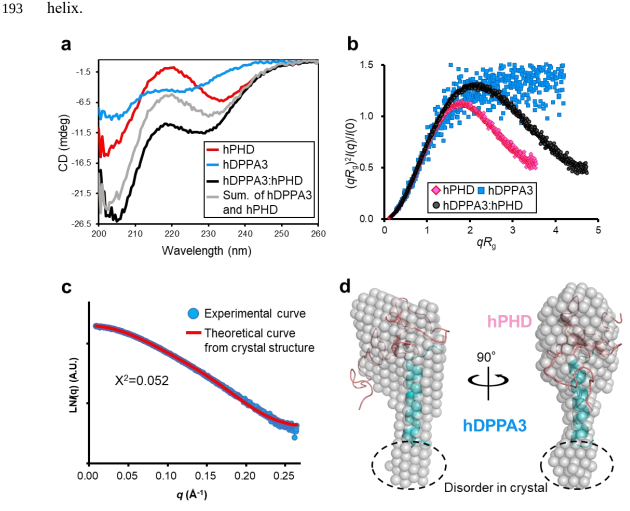
# 166 Structural feature of hPHD:hDPPA3 in solution

Intriguingly, the  $\alpha$ -helix of hDPPA3<sub>81-118</sub> has no contact with the hPHD moiety in the crystal (Figure 2a and 2c). Instead, the  $\alpha$ -helix interacts with the corresponding part of a symmetry molecule related to a crystallographic two-fold axis (Supplementary Figure 2c). This interaction in the crystal gives rise to two possibilities: the helical structure formation of hDPPA3 is an artifact of crystal packing, or the hPHD:hDPPA3 complex forms a dimer structure via the interaction mediating the  $\alpha$ -helix of hDPPA3.

Next, we examined the structure of hDPPA3<sub>81-118</sub> in solution using circular dichroism (CD) and 173 size-exclusion chromatography in line with small angle X-ray scattering (SEC-SAXS) which 174 can analyze the solution structure, oligomeric state, conformational changes and flexibility of 175 biomacromolecules at a scale ranging from a few Å to hundreds of nm (Supplementary Figure 176 3a-c, Supplementary Table 1)<sup>34</sup>. The CD spectrum exhibited that hDPPA3<sub>81-118</sub> alone showed 177 a typical random-coil spectrum (Figure 3a). The CD spectrum of hDPPA3<sub>81-118</sub> mixed with 178 hPHD showed a negative peak at 222 nm, which was lower than the sum of the spectra of 179 180 hPHD and hDPPA3<sub>81-118</sub> alone (Figure 3a), indicating that the binding of hDPPA3 to hPHD involved a coupled folding and binding mechanism. The SEC-SAXS data also supported the 181 coupled folding and binding mode of hDPPA3. The dimensionless Kratky plot showed the 182 unfolding state of sole hDPPA3<sub>81-118</sub>, whereas the hPHD:hDPPA3<sub>81-118</sub> complex was in a 183 globular state (Figure 3b). 184

SEC-SAXS experiments also revealed that the molecular mass of the measured proteins was estimated by the empirical volume of correlation  $Vc^{35}$ , resulting in a 13.0 kDa hPHD:hDDPA3<sub>81-118</sub> complex, which was highly similar to the molecular weight calculated from the amino acid sequence of the hPHD:hDPPA3<sub>81-118</sub> complex with 1:1 stoichiometry (12.2 kDa) (Supplementary Table 1). The *ab initio* model of the measured proteins showed clear results. The overall shape of the bead model was well superimposed on the crystal structure of the hPHD:hDPPA3<sub>81-118</sub> complex in the asymmetric unit (Figure 3c, 3d). These data indicated

# 192 that hDPPA3 binds to hPHD at 1:1 stoichiometry with the induction of a four-turn single $\alpha$ -



194 Figure 3: Solution structure of hDPPA3

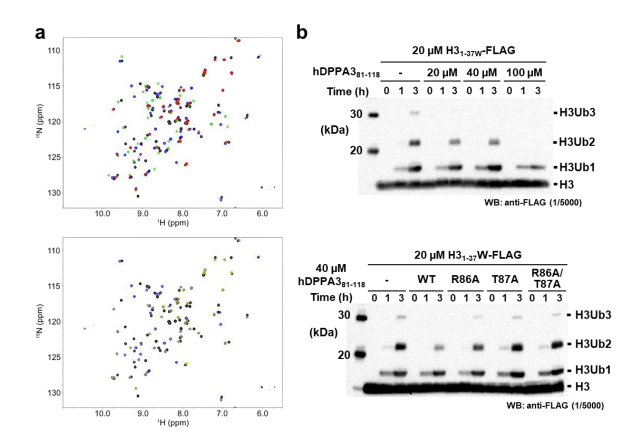
(a) CD spectra of hPHD alone (red), hDPPA3<sub>81-118</sub> alone (blue), and the hPHD in complex with 195 hDPPA3<sub>81-118</sub> (black). The sum of CD spectra of hPHD alone and hPDDA3<sub>81-118</sub> alone is shown 196 as gray. (b) Dimensionless Kratky plots of hPHD alone (red diamond), hDPPA3<sub>81-118</sub> alone 197 (blue square), and hPHD in complex with hDPPA3<sub>81-118</sub> (black circle) derived from small-angle 198 X-ray scattering (SAXS) data. (c) Structural comparison of solution and crystal structures of 199 200 the hPHD:hDPPA3<sub>81-118</sub> complex. Left shows the comparison of scattering curves derived from experimental data and the crystal structure of the hPHD:hDPPA3<sub>81-118</sub> complex. Right shows 201 202 the *ab initio* bead model of the hPHD:hDPPA3<sub>81-118</sub> complex derived from the SAXS scattering data. A low-resolution bead model (transparent gray sphere) is superimposed on the crystal 203 204 structure (cartoon). 205

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# 207 Validation of the structural data by mutational analysis

To validate our structural data and confirm the contribution of individual residues to complex 208 formation, ITC experiments were conducted using hPHD and hDPPA3 harboring mutations in 209 the VRT sequence. Mutations with deleterious effects on the interaction were R86A and T87A 210 of hDPPA3, which reduced the dissociation constant to 20.0 and 16.5 µM, respectively, and a 211 212 double mutation (R86A/T87A), which resulted in a more severe reduction in the interaction, 213 with a  $K_D$  exceeding 85.0  $\mu$ M (Figure 1c). In contrast, alanine mutations at Val85 and Leu88 in hDPPA3 had less marked effects on the hPHD:hDPPA3 interaction (Figure 1c). 214 We further investigated mutants of DPPA3 that affect dimer formation as observed in the crystal 215 structure (Supplementary Figure 4a). R98A/M102A, located at C-terminal region in the α-helix 216 of hDPPA3 and potentially interacting with hPHD of the symmetrical molecule in the crystal, 217 did not reduced the binding affinity. Similarly, M96A/L99A, which contribute to the formation 218 of the helix bundle of hDPPA3 in the crystal, also had no effect on the interaction with hPHD, 219 validating the 1:1 stoichiometry of the hPHD:hDPPA3 complex in solution. Interestingly, the 220 221 introduction of proline residue, known as a helix breaker, at both Arg93 and A97 in hDPPA3 (R93P/A97P) significantly reduced the binding affinity to hPHD, with  $K_D$  of 9.39  $\mu$ M 222 (Supplementary Figure 4a), indicating that helical structural formation following the VRT 223 motif in hDPPA3 is crucial for its interaction with hPHD. 224

Next, mutations were introduced into hPHD. Concordant with the hDPPA3 mutants, the D334A/D337A mutations in hPHD, which form an ionic-pair with Arg86 of hDPPA3, had a severe effect, reducing the binding affinity to a  $K_D$  exceeding 115 µM. The M332A hPHD mutation showed a decreased binding affinity, with a  $K_D$  of 8.07 µM (Supplementary Figure 4a). ITC data based on mutant proteins indicate that the VRT motif of hDPPA3 is important for its interaction with the UHRF1 hPHD finger.



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Figure 4: Competitive assay between hDPPA3 and the histone H3 tail.

(a) Overlay of <sup>1</sup>H-<sup>15</sup>N HSQC spectra of <sup>15</sup>N-labeled hPHD in the presence of hDPPA3<sub>81-</sub> 234 118 and/or the H3<sub>1-37W</sub> peptide at a molar ratio of 1:0:0 (black), 1:2:0 (green), 1:0:2 (blue), and 235 1:2:2 (red) of hPHD:hDPPA3:H3 (upper), and of 1:0:0 (black), 1:0:2 (blue), and 1:2:8 (yellow) 236 of hPHD:hDPPA3:H3 (lower). (b) In vitro ubiquitination assay. C-terminal FLAG tagged-H31-237 37W was ubiquitinated using in-house purified E1, E2, and human UHRF1 (E3). The 238 ubiquitinated H3 was detected using anti-FLAG antibody. Upper panel shows that 20, 40, and 239 100  $\mu$ M hDPPA3<sub>81-118</sub> was added to the reaction solution including 20  $\mu$ M of H3<sub>1-37W</sub>. The 240 lower panel presents results of an in vitro ubiquitination assay using 40 µM hDPPA3<sub>81-118</sub> 241 mutants. The gel image is representative of n = 3 independent experiments. 242

# 244 Effect of hDPPA3 on UHRF1 function

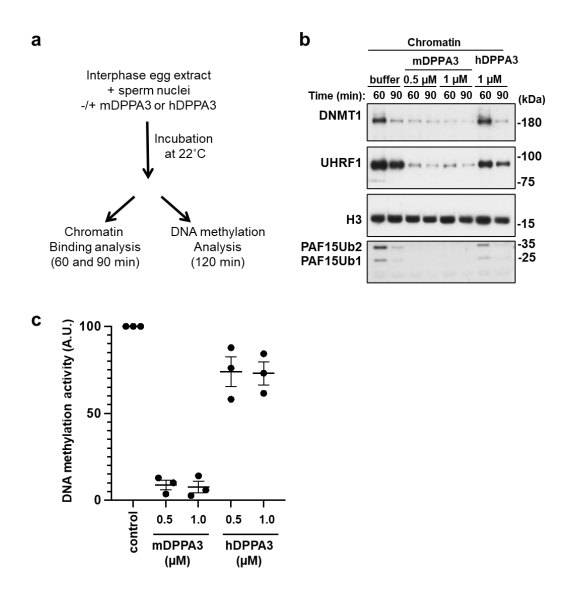
Next, to analyze whether hDPPA3 affects the biological functions of UHRF1, ubiquitination 245 of histone H3, and chromatin binding, we performed NMR titration assays and in vitro 246 biochemical experiments using recombinant proteins and Xenopus egg extracts. First, we 247 examined the competitive binding of hDPPA3 and histone H3 to hPHD because both hDPPA3 248 and histone H3 mainly bind to the acidic surface of hPHD via the <sup>85</sup>VRT<sup>87</sup> and <sup>1</sup>ART<sup>3</sup> motifs, 249 respectively. We conducted NMR titration experiments using <sup>1</sup>H-<sup>15</sup>N labeled hPHD and non-250 labeled hDPPA3<sub>81-118</sub> and/or histone H3 peptides (residues 1–37W; the H3<sub>1-37W</sub> peptide). The 251 HSQC spectrum of hPHD mixed with hDPPA3<sub>81-118</sub> and the H3<sub>1-37W</sub> peptide 252 (hPHD:hDPPA3:H3 = 1:2:2) showed most of the signals, with weakened or no intensity by the 253 broadening due to chemical exchange, suggesting that, as expected, hDPPA3<sub>81-118</sub> and the H3<sub>1-</sub> 254 <sub>37W</sub> peptide competitively bound to the acidic surface of hPHD as the shared binding site 255 (Figure 4a, upper). In the presence of excess  $H3_{1-37W}$  peptide (hPHD:hDPPA3\_{81-118}:H3 = 1:2:8), 256 hDPPA3<sub>81-118</sub> could not bind to hPHD (Figure 4a, lower). This differed from the situation with 257 258 mDPPA3, which bound to mPHD even in the presence of excess H3<sub>1-37W</sub> peptide  $(mPHD:mDPPA3:H3 = 1:2:8)^{29}$ . An *in vitro* ubiquitination assay of C-terminal FLAG-tagged 259 H3<sub>1-37W</sub> with full-length human UHRF1 also supported the weak inhibitory effect of hDPPA3. 260 hDPPA3 did not effectively inhibit ubiquitination of the histone H3 tail, whereas mDPPA3 261 showed a markedly negative effect on ubiquitination (Supplementary Figure 4b). The addition 262 of hDPPA3 to a 1-2 equimolar excess of histone H3 only slightly inhibited histone H3 263 264 ubiquitination (Figure 4b). Mutant forms of hDPPA3, which exhibited decreased binding to hPHD, failed to inhibit ubiquitination of histone H3 (Figure 4b). These findings indicate that 265 the binding of hDPPA3<sub>81-118</sub> to UHRF1 inhibits the ubiquitination activity of UHRF1 on histone 266 H3; however, the inhibitory effect was moderately weak due to the low binding affinity 267 between hDPPA3<sub>81-118</sub> and UHRF1. 268

269 Finally, we tested the ability of hDPPA3 to inhibit UHRF1 chromatin binding in *Xenopus* egg

extracts (Figure 5a). As previously reported, the addition of 0.5 µM recombinant mDPPA3 to 270 271 interphase extracts was sufficient to block UHRF1 chromatin loading, UHRF1-dependent PAF15 ubiquitylation, and DNMT1 recruitment (Figure 5b). In contrast, hDPPA3 did not 272 inhibit the chromatin binding of UHRF1 and DNMT1 recruitment, even at 1.0 µM (Figure 5b). 273 Consistently, hDPPA3 did not show significant inhibitory activity on DNA methylation in 274 *Xenopus* egg extracts compared to mDPPA3 (Figure 5c). 275 276 Taken together, the binding of hDPPA3 to hUHRF1 PHD competes with that of histone H3. However, it is noteworthy that the inhibitory effect exerted by hDPPA3 was relatively modest, 277 implying that hDPPA3 does not appear to function as a strong inhibitor of UHRF1 chromatin 278

binding, unlike mouse DPPA3.

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#### **Figure 5: Functional assay of DPPA3 using** *Xenopus* egg extracts.

(a) Experimental design for functional analysis of DPPA3 using *Xenopus* egg extracts. (b) 284 Sperm chromatin was incubated with interphase Xenopus egg extracts supplemented with 285 buffer (+buffer), FLAG-mDPPA3, or FLAG-hDPPA3. Chromatin fractions were isolated and 286 immunoblotted using the indicated antibodies. The gel image is representative of n=3287 independent experiments. (C) Sperm chromatin was added to interphase egg extracts 288 supplemented with radiolabeled S-[methyl-<sup>3</sup>H]-adenosyl-L-methionine and buffer (control), 289 FLAG-mDPPA3, or FLAG-hDPPA3. The efficiency of DNA methylation maintenance was 290 291 assessed by the incorporation of radio-labeled methyl groups from S-[methyl-<sup>3</sup>H]-adenosyl-L-292 methionine (<sup>3</sup>H-SAM) into DNA purified from the egg extracts. Data were presented as mean values  $\pm$  SD for n = 3 293

# 295 **DISCUSSION**

Our structural analysis revealed that human DPPA3 binds to hPHD solely through a conserved 296 VRT sequence motif. This finding is consistent with biochemical data showing that the binding 297 affinity of hDPPA3 to hPHD was in the sub-micro order range of  $K_D$ , with an approximately 298 30-fold decrease in the binding affinity of its mouse protein counterpart. The weak binding 299 affinity of hDPPA3 was insufficient to inhibit the chromatin binding of UHRF1 in *Xenopus* egg 300 301 extracts. Our data suggested that the inhibitory effect of hDPPA3 differs from that of mDPPA3 under similar conditions. This raises the question of whether hDPPA3 can act as an inhibitor 302 of UHRF1 in human oocytes and early embryogenesis. There are several possibilities to 303 consider in this regrd. Intrinsically disordered protein (region) containing low complexity 304 sequence frequently associates with formation of liquid-liquid phase separation  $(LLPS)^{36}$ . 305 DPPA3 Notably, sequence analysis of human and mouse using FuzDrop 306 (https://fuzdrop.bio.unipd.it) indicated that human DPPA3 exhibits a higher potential for 307 droplet formation that mouse DPPA3 (Supplementary Figure 5)<sup>37</sup>. This prediction suggests that 308 309 condensed hDPPA3 within the droplet may preferentially bind to UHRF1, thereby inhibiting the chromatin binding of UHRF1. In another situation, the level of hDPPA3 protein expression 310 in human oocytes and zygotes is key to the inhibition of UHRF1 chromatin binding. Our data 311 indicated that the binding affinity of hDPPA3 for hPHD was approximately 1.7-fold stronger 312 than that of histone H3, suggesting that a locally high concentration of hDPPA3 contributes to 313 its preferential binding to hUHRF1 to inhibit chromatin binding. Another possibility involves 314 315 post-translational modifications of histone H3. Given that the methylation of Arg2 and phosphorylation of Thr3 in histone H3 greatly impair its binding to the UHRF1 PHD finger<sup>31</sup>, 316 hDPPA3 might bind to UHRF1 even at low protein concentrations. Recently, NLRP14 317 (Nucleotide-binding oligomerization domain, leucine-rich Repeat and Pyrin domain 318 containing) has emerged as a factor related to reproduction. It interacts with UHRF1 in the 319 zygote and two-cell stages in the cytosol<sup>21,38</sup>. If cytoplasmic localization of UHRF1 is not 320

mediated by hDPPA3, it may be important for UHRF1 to interact with NLRP14 immediately after its translation into the cytoplasm. Interestingly, the cytosolic localization of the mRNA of a guanine nucleotide exchange factor, NET1, has been reported to regulate protein–protein interactions after translation, ultimately determining protein localization<sup>39</sup>.

The VRT motif in DPPA3, which binds to the acidic surface of the UHRF1 PHD finger, is well 325 conserved across various species (Figure 1b). Conversely, the amino acid sequence 326 327 corresponding to the  $\alpha$ -helix following the VRT motif showed significant diversity. Interestingly, AlphaFold2 (AF2) structural prediction indicated that mDPPA3 has both short 328 and long  $\alpha$ -helices following the VRT motif, forming an L-like shape, consistent with our NMR 329 structure of the mPHD:mDPPA3 complex (Supplementary Figure 6)<sup>40</sup>. In contrast, human 330 DPPA3 exhibits a single long  $\alpha$ -helix at the same position. AF2 predictions also suggest that 331 Homo sapience (UniProt ID: Q6W0C5), Bos taurus (A9Q1J7), Gorilla gorilla gorilla 332 (G3RB81), Saimiri boliviensis (A0A2K6SNG1), Puma concolor (A0A6P6HCW6), Nomascus 333 leucogenvs (A0A2I3H008), Crocuta crocuta (A0A6G1B388), Physeter macrocephalus 334 335 (A0A2Y9EH83), and Acinonyx jubatus (A0A6I9ZFC3) possess a single  $\alpha$ -helix, while Mus musculus (Q8QZY3), Rattus norvegicus (Q6IMK0), and Cricetulus griseus (A0A3L7H856) 336 have two  $\alpha$ -helices, consisting of both short and long  $\alpha$ -helices, as far as we could find in the 337 database (Supplementary Figure 6). These observations suggest the potential limitation of the 338 two α-helices to Rodentia and underscore the utility of AF2 structural prediction for the 339 classification of the DPPA3 function based on the helical content. The major difference in the 340 341 helical region of human and mouse DPPA3 is the substitution of a proline residue with a lysine residue at the 95th position of human DPPA3 (Figure 1a). A similar substitution is also found 342 in the species that predictably forms as single  $\alpha$ -helix. However, the K95P mutation in human 343 DPPA3 did not enhance its binding affinity for hPHD (Supplementary Figure 7a). AF2 344 prediction of the K95P mutant of hDPPA3 suggested that a single α-helix remains the 345 predominant conformation (Supplementary Figure 8), suggesting that the differences in the 346

- helical structural regions of human and mouse DPPA3 are governed by more complexmechanisms than a simple amino acid substitution.
- 349 The distinctive α-helical arrangement in hDPPA3 revealed in our structural analysis shed light
- 350 on the function of this protein in oocytes and preimplantation embryo development distinct
- 351 from the mouse DPPA3. Our results encourage further investigations into the functional
- implications of hDPPA3, potentially paving the way for novel discoveries in this context.

#### 354 Methods

#### 355 **Peptides and Primers**

- 356 The human DPPA3 peptide, residues 81-118 (NH<sup>2</sup>-<sup>81</sup>
- 357 SRRGVRTLLSVQREKMARLRYMLLGGVRTHERRPTNKE<sup>118</sup>-COOH) for
- 358 crystallography and K95P substituted hDPPA3<sub>81-118</sub> for ITC experiment were purchased from
- 359 Toray Research Center (Tokyo, Japan). Primers for site-directed mutagenesis of hDPPA3 are
- 360 listed as follows:
- 361 V85A (Forward: 5'-GAGAGGAGCAAGAACATTGCTGTGTGCA-3', Reverse: 5'-
- 362 ATGTTCTTGCTCCTCTCCTGCTCCCACCTC-3'),
- 363 R86A (Forward: 5'- AGGAGTAGCAACATTGCTGTGTGCAGAG-3', Reverse: 5'-
- 364 GCAATGTTGCTACTCCTCTCCTGCTCCCAC-3'),
- 365 T87A (Forward: 5'- AGTAAGAGCATTGCTGTCTGTGCAGAGAGA-3', Reverse: 5' 366 ACAGCAATGCTCTTACTCCTCCTGCTCC-3'),
- 367 L88A (Forward: 5'- AAGAACAGCGCTGTCTGTGCAGAGAGAAAA-3', Reverse: 5'-
- 368 CAGACAGCGCTGTTCTTACTCCTCTCCTGC-3')
- 369 R86A/T87A (Forward: 5'- AGGAGTAGCAGCATTGCTGTGTGCAGAG-3', Reverse: 5'-
- 370 ACAGCAATGCTGCTACTCCTCTCCTGCTCC-3').
- 371 R98A/M102A (Forward: 5'- GGCAGCATTGAGATACGCGTTACTCGGCGGAGTTC -3',
- 372 Reverse: 5'- GTAACGCGTATCTCAATGCTGCCATCTTTTCTCTC -3').
- 373 M96A/L99A (Forward: 5'- AAAGGCGGCAAGAGCGAGATACATGTTACTCGGCG -3',
- 374 Reverse: 5'- ATCTCGCTCTTGCCGCCTTTTCTCTCTGCACAGAC -3').
- 375 R93P/A97P (Forward: 5'- GCAGCCAGAAAAGATGCCAAGATTGAGATACATGT -3',
- 376 Reverse: 5'- ATCTTGGCATCTTTTCTGGCTGCACAGACAGCAAT -3').
- 377
- 378 **Protein expression and purification**
- Human UHRF1 PHD finger (residues 299-366) for crystallography, SAXS, NMR, CD, and

ITC experiments was expressed in *Escherichia coli* (*E.coli*) and purified according to previous 380 paper<sup>31</sup>. Briefly, hPHD was expressed as a GST-fusion protein and purified using glutathione 381 Sepharose 4B (GS4B), anion exchange (HiTrap Q) and 26/600 Supedex 75 chromatography 382 (Cytiva). hDPPA3, residues 81–118, for SAXS, NMR, CD, ITC and ubiquitination experiments 383 was expressed as a six histidine-tagged ubiquitin (His-Ub) fusion protein. The protein was 384 expressed in E. coli BL21 (DE3) in Luria-Bertani medium (LB) containing 12.5 µg/ml 385 kanamycin. When the optical density at 660 nm (O.D.660) of the cells reached 0.7, 0.4 mM 386 isopropyl  $\beta$ -d-thiogalactoside (IPTG) was added to the medium and the cells were further 387 harvested for 6 h at 30 °C. The cells were suspended in lysis buffer (40 mM Tris-HCl [pH7.5], 388 400 mM NaCl and 30 mM imidazole). After cell lysis by sonication and removal of cell debris 389 by centrifugation, the supernatant was loaded onto a histidine-tag affinity column Ni Sepharose 390 391 6 Fast Flow (Cytiva), and the sample was eluted from the column using elution buffer containing 500 mM imidazole. Next, the His-Ub tag was removed by Saccharomyces 392 cerevisiae ubiquitin carboxyl-terminal hydrolase YUH1. The sample was further purified using 393 394 HiTrap SP HP cation-exchange chromatography (Cytiva) and finally purified using HiLoad 26/600 Superdex 75 size-exclusion chromatography equilibrated with  $1 \times ITC$  buffer (10 mM 395 HEPES (pH7.5), 150 mM NaCl, 0.25 mM tris (2-carboxyethyl)phosphine (TCEP)). The H3 396 peptide (residues 1–36 with an additional tryptophan residue at their C-terminus, hereafter H3<sub>1</sub>-397 <sub>37W</sub>), mouse DPPA3, full-length mouse UHRF1, full-length human UHRF1, mouse UBA1 and 398 human UBE2D3 for the *in vitro* ubiquitination assay were purified according to previous 399 reports<sup>17,29</sup>. 400

For the preparation of <sup>15</sup>N-labeled or <sup>13</sup>C,<sup>15</sup>N-double labeled hPHD, M9 minimal media containing 0.5 g/l <sup>15</sup>NH<sub>4</sub>Cl or 0.5 g/l <sup>15</sup>NH<sub>4</sub>Cl and 1 g/l <sup>13</sup>C-glucose was used instead of LB media. Site-directed mutagenesis of hPHD and hDPPA3<sub>81-118</sub> was performed by designing two primers containing the mutations. The mutants of hDPPA3<sub>81-118</sub> and the labeled hPHD were purified using the same protocol. The mutants of hDPPA3<sub>81-118</sub> and the labeled hPHD were

406 purified by the same protocol.

407

# 408 Crystallography of hPHD in complex with hDPPA3 peptide

The hPHD:hDPPA3<sub>81-118</sub> complex was prepared by adding an equi-molar excess of hDPPA3 409 peptide to hPHD prior to crystallization. The crystal was obtained using an 8 mg/ml 410 concentration of the complex at 4°C and the hanging drop vapor diffusion method with a 411 412 reservoir solution containing 100 mM Tris-HCl (pH 8.5) and 2 M Ammonium sulfate. The crystal was directly frozen in liquid nitrogen using a cryoprotectant containing 25% (v/v) 413 ethylene glycol. The X-ray diffraction data were collected at a wavelength of 0.98000 Å on a 414 Pilatus3 6M detector in beam line BL-17A at Photon Factory (Tsukuba, Japan) and scaled at 415 2.40 Å resolution using the program XDS package<sup>41</sup> and Aimless<sup>42</sup>. After molecular 416 replacement by PHASER<sup>43</sup> using human PHD finger (PDB: 3ASL) as a search model and 417 several cycles of model refinement by PHENIX<sup>44</sup>, the final model converged at 2.40 Å 418 resolution with a crystallographic *R*-factor of 23.3% and a free *R*-factor of 26.6%. 419

The crystallographic data and refinement statistics are given in Table 1. Figures were generated
using PyMol (<u>http://www.pymol.org</u>).

422

#### 423 **NMR**

All NMR experiments were performed using a Bruker BioSpin Avance III HD spectrometers 424 with TCI triple-resonance cryogenic probe-heads and basic <sup>1</sup>H resonance frequency of 600.03 425 426 and 800.23 MHz. Three-dimensional (3D) spectra for backbone signal assignments, including HNCACB, CACB(CO)NH, HNCA, HN(CO)CA, HNCO, and HN(CA)CO, were acquired at 427 293 K for 520 µM [<sup>13</sup>C, <sup>15</sup>N]-hPHD dissolved in PBS buffer (pH 7.0) containing 1 mM DTT 428 and 5% D<sub>2</sub>O. For the complex state, 260 µM [<sup>13</sup>C, <sup>15</sup>N]-hPHD with hDPPA3<sub>811-118</sub> at molar 429 ratio of 1:2 was used in the buffer same as the free state. The spectral widths (total number of 430 data points) of each spectrum were 18 ppm (2048) for the <sup>1</sup>H dimension and 24 ppm (192) for 431

432	the <sup>15</sup> N dimension. All 3D spectra were acquired using non-uniform sampling (NUS) to
433	randomly reduce the $t_1$ and $t_2$ time-domain data points by 25%. The uniformly sampled data
434	were reconstructed from the raw NMR data using various techniques such as IST or SMILE <sup>45,46</sup>
435	All NMR spectra were processed using NMRPipe <sup>47</sup> . For NMR analysis, an integrated package
436	of NMR tools named MagRO-NMRViewJ, version 2.01.41 <sup>48</sup> , on NMRView was used <sup>49</sup> .
437	For the competition experiments, $^1\text{H-}^{15}\text{N}$ HSQC spectra were measured at 293K for 60 $\mu\text{M}$
438	[ <sup>15</sup> N]-hPHD in the presence of hDPPA <sub>81-118</sub> and/or the H3 <sub>1-37W</sub> peptide at molar ratios
439	(hPHD:hPDDA3:H3) of 1:0:0, 1:2:0, 1:0:2, 1:2:2 and 1:2:8.
440	
441	ITC measurements
442	Microcal PEAQ-ITC (Malvern) was used for ITC measurements. Wild-type and mutants of
443	hPHD and hDPPA3 were dissolved in ITC buffer (10 mM HEPES [pH 7.5] buffer containing
444	150 mM NaCl and 0.25 mM TCEP). All measurements were carried out at 293 K. The data
445	were analyzed with Microcal PEAQ-ITC analysis software using a one-site model. For each
446	interaction, at least three independent titration experiments were performed to show the

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Far-UV circular dichroism (CD) spectra were obtained using a JASCO J-1100 model spectrometer. All samples were prepared at a concentration of 20  $\mu$ M, dissolved in 10 mM HEPES [pH7.5] buffer containing 150 mM NaCl, 0.25 mM TCEP. The measurements were performed at 293 K with a path length of 1 mm.

dissociation constants with mean standard deviation.

454

# 455 SEC-SAXS

456 SAXS data were collected on Photon Factory BL-10C using an HPLC Nexera/Prominence-I

457 (Shimazu) integrated SAXS set-up<sup>50</sup>. 50 μL of 12 mg/mL hPHD and hPHD:hDPPA3<sub>81-118</sub>

<sup>449</sup> **CD** 

complex and 20 mg/ml hDPPA3<sub>81-118</sub> were loaded onto a Superdex® 200 Increase 5/150 GL 458 (Cytiva) pre-equilibrated with 20 mM Tris-HCl (pH 7.5), 150 mM NaCl, 2 mM DTT, 10 µM 459 zinc acetate and 5% glycerol at a flow rate of 0.25 mL/min at 20 °C. The flow rate was reduced 460 to 0.025 mL/min at an elution volume of 1.9-2.8 mL. X-ray scattering data were collected 461 every 20 s on a PILATUS3 2 M detector over an angular range of  $q_{\min} = 0.00690 \text{ Å}^{-1}$  to 462  $q_{\text{max}} = 0.27815 \text{ Å}^{-1}$ . The UV spectra at the range of 200 to 450 nm were recorded every 10 s. 463 Circular averaging and buffer subtraction were carried out using the program SAngler<sup>51</sup> to 464 obtain one-dimensional scattering data I(q) as a function of q  $(q = 4\pi \sin\theta/\lambda)$ , where  $2\theta$  is the 465 scattering angle and  $\lambda$  is the X-ray wavelength 1.5 Å). The scattering intensity was normalized 466 on an absolute scale using the scattering intensity of water<sup>42</sup>. The multiple concentrations of 467 the scattering data around the peak at A280, namely the ascending and descending parts of the 468 chromatography peak, and I(0) were extrapolated to zero-concentration using MOLASS<sup>52</sup>. The 469 molecular mass of the measured proteins was estimated using the empirical volume of 470 correlation,  $V_c$ , showing no aggregation of the measured sample<sup>35</sup>. The radius of gyration  $R_g$ 471 and forward scattering intensity I(0) were estimated from the Guinier plot of I(q) in the smaller-472 angle region of  $qR_g < 1.3$ . The distance distribution function, P(r), was calculated using the 473 program GNOM<sup>53</sup>. The maximum particle dimension  $D_{\text{max}}$  was estimated from the P(r)474 function as the distance r for which P(r) = 0. The scattering profile of the crystal structure of 475 hPHD:hDPPA3<sub>81-118</sub> was computed using CRYSOL<sup>54</sup> software. Ab initio model of 476 hPHD:hDPPA3<sub>81-118</sub> was created using GASBOR and DAMAVER<sup>55,56</sup> 477

478

# 479 *In vitro* ubiquitination assay

480 Protein expression in *E. coli* and purification of mouse UBA1 (E1), human UBE2D3 (E2), 481 human UHRF1 (E3), C-terminal FLAG tagged-H3<sub>1-37W</sub> and ubiquitin were performed 482 according to previous reports<sup>17</sup>. The ubiquitination reaction mixtures contained 100  $\mu$ M 483 ubiquitin, 200 nM E1, 8  $\mu$ M E2, 3  $\mu$ M E3, 5 mM ATP, and 20  $\mu$ M C-terminal FLAG tagged-

H3<sub>1-37W</sub> in the presence and absence of hDPPA3<sub>81-118</sub> in ubiquitination reaction buffer (50 mM
Tris-HCl [pH 8.0], 50 mM NaCl, 5 mM MgCl<sub>2</sub>, 0.1% Triton X-100, 2 mM DTT). The mixture
was incubated at 30°C for 3 h and the reaction was stopped by adding 3×SDS loading buffer.
The reaction was analyzed by SDS-PAGE, followed by Western blotting using a 1/5,000 diluted
anti-FLAG antibody (Cell Signaling Technology, #2368).

489

# 490 *Xenopus* interphase egg extracts and purification of chromatin

Xenopus laevis was purchased from Kato-S Kagaku and handled according to animal care 491 regulations at the University of Tokyo. Interphase egg extracts were prepared as described 492 previously<sup>12</sup>. Unfertilized *Xenopus* eggs were dejellied in 2.5% thioglycolic acid-NaOH (pH 493 8.2) and washed three times in 0.2×MMR buffer (5 mM HEPES-KOH [pH 7.6], 100 mM NaCl, 494 495 2 mM KCl, 0.1 mM EDTA, 1 mM MgCl<sub>2</sub>, 2 mM CaCl<sub>2</sub>). After activation in 1×MMR supplemented with 0.3 µg/mL calcium ionophore, eggs were washed four times with EB buffer 496 (10 mM HEPES-KOH [pH 7.7], 100 mM KCl, 0.1 mM CaCl<sub>2</sub>, 1 mM MgCl<sub>2</sub>, 50 mM sucrose). 497 498 Packed eggs were crushed by centrifugation (BECKMAN, Avanti J-E, JS13.1 swinging rotor) for 20 min at 18,973×g. Egg extracts were supplemented with 50  $\mu$ g/mL cycloheximide, 20 499 µg/mL cytochalasin B, 1 mM DTT, 2 µg/mL aprotinin, and 50 µg/mL leupeptin and clarified 500 for 20 min at 48,400×g (Hitachi, CP100NX, P55ST2 swinging rotor). The cytoplasmic extracts 501 were aliquoted and stored at -80°C. Chromatin purification after incubation in egg extracts was 502 performed as previously described with modifications. Sperm nuclei were incubated in egg 503 504 extracts supplemented with an ATP regeneration system (20 mM phosphocreatine, 4 mM ATP, and 5 µg/mL creatine phosphokinase) at 3000–4000 nuclei/µL at 22°C. Aliquots (15 µL) were 505 diluted with 150-200 µL of chromatin purification buffer (CPB; 50 mM KCl, 5 mM MgCl<sub>2</sub>, 20 506 mM HEPES-KOH [pH 7.7]) containing 0.1% NP-40, 2% sucrose, and 2 mM NEM. After 507 incubation on ice for 5 min, the extracts were layered over 1.5 mL CPB containing 30% sucrose 508 and centrifuged at 15,000×g for 10 min at 4°C. Chromatin pellets were resuspended in 509

1×Laemmli sample buffer, heated for 5 min and analyzed by SDS-PAGE. Recombinant FLAG-510 tagged mDPPA3 and hDPPA3 were added to egg extracts at 0.5-1.0 µM. For FLAG-tagged 511 protein expression in insect cells, 3×FLAG-tagged *mDppa3* or *hDppa3* were sub-cloned into 512 pVL1392 vector. Baculoviruses were produced using a BD BaculoGold Transfection Kit and a 513 BestBac Transfection Kit (BD Biosciences) following the manufacturer's protocol. Proteins 514 515 were expressed in Sf9 insect cells by infection with viruses expressing 3×FLAG-tagged 516 mDPPA3 WT or its mutants for 72 h at 27 °C. Sf9 cells from a 750 ml culture were collected and lysed by resuspending them in 30 ml lysis buffer, followed by incubation on ice for 10 min. 517 The soluble fraction was obtained after centrifugation of the lysate at 15,000×g for 15 min at 518 4 °C. The soluble fraction was incubated with 250 µL anti-FLAG M2 affinity resin equilibrated 519 with lysis buffer for 4 h at 4 °C. The beads were collected and washed with 10 ml wash buffer 520 and then with 5 ml of EB [20 mM HEPES-KOH (pH 7.5), 100 mM KCl, 5 mM MgCl<sub>2</sub>] 521 containing 1 mM DTT. Each recombinant protein was eluted twice in 250 µL EB containing 1 522 mM DTT and 250 µg/ml 3×FLAG peptide (Sigma-Aldrich). The eluates were pooled and 523 524 concentrated using a Vivaspin 500 (GE Healthcare).

# 525 Data Availability

- 526 Coordinate of atomic model of human UHRF1 PHD finger in complex with human DPPA3
- 527 was deposited in the Protein Data Bank with accession code 8WMS. All data needed to evaluate
- 528 the conclusions in the paper are presented in the paper and/or Supplementary Materials.
- 529 Additional data related to this paper may be requested from the authors.

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#### 661 Author Contributions

K.A. conceived the study and experimental design, analyzed the experiments, and wrote the manuscript. N.S. and K.A. performed the protein purification, X-ray crystallography, SAXS measurements, and ITC experiments. N.S. performed CD experiments. N.S. and N.N. performed *in vitro* ubiquitination assays. S.H. and T. K. performed NMR experiments and analyzed the data. Y.C. IM. H. M. N. and A.N. performed the biochemical assays using *Xenopus* egg extracts.

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# 676 **CONFLICT OF INTEREST**

The authors declare no competing interests.

#### 679 Table 1. Data collection and refinement statistics

	hPHD:hDPPA3 680	
	(PDB:8WMS)	
Data collection		
Space group	<i>I</i> 4 <sub>1</sub> 22	
Cell dimensions		
<i>a</i> , <i>b</i> , <i>c</i> (Å)	77.80 77.80 140.67	
Resolution (Å)	43.33-2.40 (2.49-2.40)*	
$R_{\rm meas}$ (%)	7.0 (53.7)*	
<i>R</i> <sub>pim</sub> (%)	2.6 (21.0)*	
Mean $(I/\sigma(I))$	12.5 (2.5)*	
$CC_{1/2}$	99.9 (89.6)*	
Completeness (%)	99.7 (99.6)*	
Redundancy	5.4 (5.7)*	
Total reflections	47,813 (5,172)*	
Unique reflections	8,775 (903)*	
Refinement		
Resolution (Å)	43.33-2.40	
No. reflections	8,693	
$R_{\rm work} / R_{\rm free}$ (%)	23.3 / 26.6	
No. atoms		
hPHD	514	
hDPPA3	190	
zinc	4	
<i>B</i> factors (Å <sup>2</sup> )		
hPHD	79.9	
hDPPA3	83.5	
zinc	67.6	
R.m.s. deviations		
Bond lengths (Å)	0.004	
Bond angles (°)	0.803	

\* ( ) Values in parentheses are for the highest-resolution shell.