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Poly ADP-ribose signaling is dysregulated in Huntington disease

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Affiliations are included on p. 10.

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Huntington disease (HD) is a genetic neurodegenerative disease caused by cytosine, adenine, guanine (CAG) expansion in the Huntingtin (HTT) gene, translating to an expanded polyglutamine tract in the HTT protein. Age at disease onset correlates to CAG repeat length but varies by decades between individuals with identical repeat lengths. Genome-wide association studies link HD modification to DNA repair and mitochondrial health pathways. Clinical studies show elevated DNA damage in HD, even at the premanifest stage. A major DNA repair node influencing neurodegenerative disease is the PARP pathway. Accumulation of poly adenosine diphosphate (ADP)-ribose (PAR) has been implicated in Alzheimer and Parkinson diseases, as well as cerebellar ataxia. We report that HD mutation carriers have lower cerebrospinal fluid PAR levels than healthy controls, starting at the premanifest stage. Human HD induced pluripotent stem cell-derived neurons and patient-derived fibroblasts have diminished PAR response in the context of elevated DNA damage. We have defined a PAR-binding motif in HTT, detected HTT complexed with PARylated proteins in human cells during stress, and localized HTT to mitotic chromosomes upon inhibition of PAR degradation. Direct HTT PAR binding was measured by fluorescence polarization and visualized by atomic force microscopy at the single molecule level. While wild-type and mutant HTT did not differ in their PAR binding ability, purified wild-type HTT protein increased in vitro PARP1 activity while mutant HTT did not. These results provide insight into an early molecular mechanism of HD, suggesting possible targets for the design of early preventive therapies.

PARylation | Huntington's disease | huntingtin | PARP1 | poly ADP-ribose

Huntington disease (HD) is an autosomal dominant genetic neurodegenerative disease caused by a cytosine, adenine, guanine (CAG) expansion in exon1 of the *Huntingtin* (*HTT*) gene, which translates to an expanded polyglutamine tract in the HTT protein. HD is characterized by psychiatric, cognitive, and motor disturbances for which some symptom management treatments, but no disease-modifying therapies, exist. Age at disease onset is negatively correlated with CAG expansion length, and onset typically occurs in the third or fourth decades of life (1). However, age at disease onset in individuals with the same CAG length can vary by decades (2, 3). This variability has an inherited component (4, 5), suggesting that other genes may act as modifiers of disease onset. Genome-wide association studies have primarily implicated DNA repair and maintenance pathways as such modifiers (2, 3, 6, 7). While the pathogenic mechanisms have yet to be elucidated, single-nucleotide polymorphisms in DNA repair genes that affect somatic instability of the HTT locus have been linked to age at onset (8–10), and elevated levels of DNA damage in HD patient-derived samples have been reported (11–22).

The connection between DNA repair and neurodegeneration is not unique to HD and has been described for Alzheimer disease (23), Parkinson disease (24), spinocerebellar ataxias (25, 26), and amyotrophic lateral sclerosis (ALS) (27, 28). Inherited mutations in a number of DNA repair genes cause neurodegenerative disorders (29–33), and of the nine neurodegenerative genetic polyglutamine diseases, seven of the causative proteins have been implicated in the DNA damage response (34). HTT forms a transcription-coupled DNA repair complex with RNA polymerase II subunit A, polynucleotide kinase-phosphatase, and ataxin-3, and mutant HTT interferes with this function (21, 22). We have previously reported that HTT interacts with ataxia-telangiectasia mutated (ATM), and that ATM kinase activity is required for HTT–chromatin interaction in response to oxidative stress (12). We have more recently identified a similar relationship between ATM and ataxin-1, the protein product of the *ATXN1* gene mutated in spinocerebellar ataxia 1 (35). ATM signaling is also dysregulated in HD models and brain tissue (11), placing it as a DNA damage repair node in neurodegenerative diseases.

Significance

A consensus on dysfunctional DNA repair has emerged in neurodegenerative disease research, with elevated poly adenosine diphosphate (ADP)ribose (PAR) signaling more recently implicated. In contrast, we have identified a deficient PAR response in Huntington disease (HD) patient spinal fluid samples and cells. This may be explained by the inability of huntingtin protein bearing the HD-causing mutation to stimulate production of PAR the way the wild-type protein does. Since drugs that target PAR production and degradation have already been developed, these findings present an exciting avenue for therapeutic intervention for HD.

Competing interest statement: E.J.W. reports consultancy/ advisory board memberships with Annexon, Remix Therapeutics, Hoffman La Roche Ltd., Ionis Pharmaceuticals, PTC Therapeutics, Takeda, Teitur Trophics, Triplet Therapeutics and Vico Therapeutics. All honoraria for these consultancies were paid through the offices of UCL Consultants Ltd., a wholly owned subsidiary of UNIversity College London. F.B.R. is a Medpace UK Ltd. employee and was a University College London employee during the conduct of this study. F.B.R. has provided consultancy services to GLG and F. Hoffmann-La Roche Ltd. L.M.B. currently holds consultancy contracts with Annexon Biosciences, Remix Therapeutics, and LoQus23 Therapeutics Ltd. via UCL Consultants Ltd. R.T. has past consultancy with Novartis AG, PTC Therapeutics and Mitokinin LLC.

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Another such node of DNA damage repair is PARP signaling, which has been implicated in Alzheimer, Parkinson, and HDs (reviewed in ref. 36) as well as progressive cerebellar ataxia (32) and ALS (28). Of the 17 members of the PARP family, PARP1 and PARP2 are critical in the DNA damage response, with PARP1 accounting for 80 to 90% of DNA repair-related activity (37, 38). Upon activation by DNA breaks, PARP1 and PARP2 use adenosine diphosphate (ADP)-ribose units from nicotinamide adenine dinucleotide (NAD⁺) as building blocks to generate poly ADP-ribose (PAR) chains of varying length and branching structure. PAR chains act as recruitment scaffolds for DNA repair proteins (39–41). Part of the repair process involves PAR degradation by PAR glycohydrolase (PARG), which is required to allow DNA repair proteins access to the damaged DNA (42), and to recycle ATP precursor metabolites (43, 44).

Excessive PAR polymerization, or lack of degradation, results in NAD⁺ depletion, energy crisis, and cell death by necrosis (45). Overactivation of PARP1 can also trigger the nonapoptotic programmed cell death termed parthanatos (46), which is particularly important in neurodegenerative disease (47). Hyper-PARylation has been implicated in several neurodegenerative diseases (32, 48–50), including HD (51–53).

Here, we show that PAR signaling is dysregulated in HD patient cells and cerebrospinal fluid (CSF). Analysis of HD mutation carrier CSF samples revealed lower total PAR levels than control samples, a difference evident from the premanifest stage. Elevated DNA damage levels in HD induced pluripotent stem cell (iPSC)-derived neurons and patient-derived fibroblasts were not reflected by increased PAR levels, and HD patient-derived cells had lower PARP1/2 inhibitor IC₅₀ than control cells. Consistent with this, wild-type purified recombinant HTT increased autoPARylation of purified recombinant PARP1 in an in vitro assay, while mutant HTT was deficient in this capacity. We detected endogenous HTT complexed with PARylated proteins in human retinal epithelial cells, which prompted analysis of the HTT structure and identification of a PAR-binding motif (PBM). Fluorescence polarization and atomic force microscopy revealed a direct interaction between HTT protein and PAR polymers. While PAR binding by wild-type or mutant HTT did not differ, mutant HTT was not able to stimulate PARP1 activity while wild-type protein demonstrated this activity. This provides a possible mechanistic link to the observations made in patient cells and CSF. In addition to uncovering a role for the normal HTT protein in PAR biology, these results provide insight into the very early molecular mechanism of HD pathogenesis, to potentially reveal targets for an early preventive therapy for HD.

Results

The PAR Response Is Deficient in HD Patient-Derived Samples.

Elevated PAR levels have been observed in human samples from multiple neurodegenerative diseases (32, 48–50). Although PARP1 inhibition was reported to be neuroprotective in a mouse model of HD (51, 52), and elevated PARP1 levels were observed in the neurons and glia of the caudate nucleus from HD patients (53), it remains unknown whether hyper-PARylation contributes to HD pathology. As cancer drugs targeting PARP1 are under consideration for repurposing to treat neurodegenerative diseases (54, 55), investigation of their potential for HD is imperative. To test patient-relevant CNS-specific samples, we measured PAR levels in CSF from HD patients compared to controls. Samples were obtained from the HD-CSF study, an 80-participant crosssectional study of HD mutation carriers and matched healthy controls (56, 57) (see *Materials and Methods* section for details). In contrast to similar analyses in Parkinson's disease (50), we found that PAR levels were significantly lower in the CSF from both premanifest (P = 0.0001) and manifest HD patients (P = 0.0004) compared to controls, and that gene status had a large effect on PAR levels in CSF ($\eta_p^2 = 0.35$) (Fig. 1). PAR levels in the CSF of HD mutation carriers did not correlate with CSF mutant HTT or neurofilament light chain (NfL) levels, both of which were shown to correlate with disease severity in HD patients (56, 57) (*SI Appendix*, Fig. S1). In addition, CSF PAR levels in HD mutation carriers did not correlate with clinical measures of disease progression such as disease burden score, total functional capacity, or total motor score (*SI Appendix*, Fig. S2). Yet, lower PAR levels correlated with the presence of the mutant HTT allele.

The reduced PAR levels in the CSF of premanifest and manifest HD patients were unexpected given the numerous reports of increased DNA damage in HD patient samples and models. For example, iPSC-derived neurons bearing the HD mutation have increased DNA breaks and expression of damage markers phospho-ATM, γ H2AX, and phospho-p53 (21, 22). We therefore aimed to determine whether this increase in DNA damage was reflected by elevated PAR levels. As shown in Fig. 2*A*, we found no corresponding increase in PAR levels in HD iPSC-derived neurons. In fact, neurons bearing a juvenile-onset HD allele (Q77) had lower PAR levels than control, consistent with the observations made in patient CSF.

We and others have previously reported elevated levels of DNA damage in fibroblasts derived from HD patients compared to controls (12, 18, 20). We therefore measured DNA damage and PAR levels in control cells bearing wild-type alleles that encode 21 or 18 poly-glutamine residues (TruHD-Q21Q18), and HD patient-derived cells carrying one expanded allele (TruHD-Q43Q17), or two expanded alleles (TruHD-Q50Q40), TruHD cells (58). We employed Repair Assisted Damage Detection (RADD) (59) to measure DNA breaks and PAR levels in the same cells. RADD directly detects DNA damage using DNA processing enzymes to detect and modify sites of DNA damage for a subsequent gap-filling fluorescent labeling reaction. Like HD iPSC-derived neurons, human telomerase reverse transcriptase (hTERT)-immortalized HD patient-derived fibroblasts



Fig. 1. PAR levels are reduced in premanifest and manifest HD patient CSF. CSF samples from control, premanifest HD, and manifest HD subjects were blinded and analyzed for PAR levels by ELISA. Group comparisons were assessed using multiple regression with post hoc Wald tests. *Survives Bonferroni correction for multiple comparisons.



Fig. 2. The PAR response is deficient in HD cells. (*A*) iPSC-derived neurons were fixed and stained with neuronal marker Map2 (red) and MABE1031 PAR detection reagent (green). Nuclear PAR intensity in Map2-positive cells was measured using CellProfiler. Data from six (CTR Q18 and HD Q53) or four (JHD Q77) differentiation replicates are shown (n = 100 to 300 nuclei per cell line). Results were analyzed by the Kruskal–Wallis test and corrected for multiple comparisons using Dunn's test. Error bars: SEM. (*B*) hTERT-immortalized fibroblasts from healthy control (TruHD-Q21Q18) and HD patients (TruHD-Q43Q17 and TruHD-Q50Q40) were treated with 100 mM KBrO₃ for 30 min followed by Repair Assisted Damage Detection (RADD) to detect DNA damage, and costaining with MABE1031 PAR detection reagent. Representative images of TruHD-Q21Q18 cells are shown. Nuclear RADD and PAR intensity were measured using CellProfiler, mean intensity recorded for each image (18 images per condition; >500 cells), and values normalized to the control condition. Data from three independent experiments are shown. Results were analyzed by two-way ANOVA and corrected for multiple comparisons using Tukey's test. Error bars: SEM. (Scale bars: 50 microns.)

did not exhibit the increase in PAR levels expected to occur in the context of elevated DNA damage (Fig. 2*B*). This was the case under conditions of oxidative stress, which we have previously shown to be associated with elevated DNA damage in these cells by comet assay (12), as well as under basal conditions, in which increased damage was also detected by RADD (Fig. 2*B*). To ensure that the PAR detection assay is within the dynamic range and the signal has not reached saturation, we quantified PAR signal intensity in a KBrO₃ doseresponse experiment. As shown in *SI Appendix*, Fig. S3, a treatment of 100 mM KBrO₃ for 30 min produces a PAR response within the dynamic range. Further, HD cells exhibited lower PAR levels than wild-type cells at the other KBrO₃ doses.

Since hTERT expression is equal across the immortalized fibroblast lines (58), and similar results were obtained in iPSC-derived neurons and patient CSF, we did not anticipate a confounding effect of hTERT overexpression. This was confirmed by repeating the experiment in the primary fibroblasts from which the TruHD lines were derived (*SI Appendix*, Fig. S4). It should be noted that although we did not observe the expected elevated PAR phenotype in HD patient-derived cells, PAR was generated in response to oxidative stress, indicating that they are not fully deficient in this capacity (Fig. 2B). To further characterize this PAR signaling deficiency, we correlated per-nucleus RADD and PAR signal intensities for control and HD patient-derived cells and found them to be correlated (*SI Appendix*, Fig. S5). As expected, the correlation increased with KBrO₃ treatment; however, HD cells had lower Spearman r values than wild-type cells under both conditions. This suggests that the PAR response is not fully deficient but is subdued in the HD context, as would be expected for a late onset, slowly progressing disease. Together, the reduced PAR levels in HD patient CSF, and lack of elevated PAR in the context of elevated DNA damage in HD neurons and patient-derived fibroblasts, suggest that the PAR response is deficient in HD patients.

We next examined the mechanisms of PAR production and degradation in TruHD cells. During the DNA repair process, PAR is rapidly generated by PARP1 and PARP2, and degraded by enzymes such as PARG. To determine whether the lower-than-expected PAR levels were due to increased PARG activity, we performed dose-response experiments to measure the PARG inhibitor PDD00017273 IC_{50} (60) and found no difference between HD and control cells (Fig. 3A). In contrast, dose-response experiments with the PARP1/2 inhibitor veliparib revealed lower veliparib IC₅₀ in HD cells compared to controls (TruHD-Q21Q18 IC₅₀ = 140 nM; TruHD-Q43Q17 IC₅₀ = 60 nM; TruHD-Q50Q40 IC₅₀ = 60 nM) (Fig. 3B). This is despite similar levels of PARP1, PARP2, and PARG across cell lines (SI Appendix, Fig. S6A) and equal PARP1 chromatin retention upon oxidative stress (SI Appendix, Fig. S6B). To ensure that the lowered IC_{50} is not an artifact specific to veliparib, we repeated the experiment with talazoparib, which acts by a different mechanism than veliparib (61, 62), and found a reduced IC_{50} in HD cells (SI Appendix, Fig S6C), as was seen with veliparib. To address whether the reduced PARP inhibitor IC₅₀ values were a direct effect of HTT, we tested the autoPARylation activity of purified recombinant PARP1 in the presence of purified recombinant HTT-HAP40 Q23 or HTT-HAP40 Q54. As shown in Fig. 3*C*, wild-type HTT stimulated PARP1 activity in a dose-dependent manner, while mutant HTT had no effect. This provides a possible mechanistic link to the reduced PARP inhibitor IC_{50} values, and subdued PAR response observed in HD patient-derived fibroblasts, as well as the reduced PAR levels in HD iPSC-derived neurons and CSF from HD patients.

HTT Interacts with PARylated Proteins. Mutations associated with progressive cerebellar ataxia with oculomotor apraxia type 1 result in reduced expression of the scaffolding protein XRCC1, causing persistent unrepaired DNA damage and concomitant prolonged PARP1 activity (32). Since PAR binding is important for the scaffolding function of XRCC1 in DNA repair complex formation (63), we sought to determine whether HTT could also bind PAR.

We first interrogated the results of a mass spectrometry study identifying proteins that interact with HTT under conditions of oxidative



Fig. 3. PARP1/2 activity is higher in the presence of wild-type HTT. Fibroblasts were pretreated with increasing doses of PDD00017273 (*A*) or veliparib (*B*) for 30 min followed by 100 mM KBrO3 for 30 min in the presence of inhibitor. Veliparib dose-response was carried out in the presence of 5 μ M PARG inhibitor to enable pan-ADP-ribose detection by MABE1016. EC₅₀ and IC₅₀ values were calculated from nuclear PAR staining intensity (10 to 12 images per condition; >800 cells) using GraphPad Prism. Error bars = SEM for four (PARG inhibitor) or eight (veliparib) experiments. ****P* < 0.0001 (Brown–Forsythe and Welch ANOVA tests). (*C*) 10 fmol recombinant PARP1 was incubated with the indicated amounts of recombinant HTT-HAP40 for 2 h at 30 °C. Reactions were separated by SDS-PAGE and immunoblotted with MABE1016 pan-ADP-ribose detection reagent. Signal intensities were quantified using ImageJ. Values from three (HTT-HAP40 Q54) or four (HTT-HAP40 Q23) experiments are shown.

stress (Dataset S1). The list of HTT interactors was compared to a compiled list of PARylated proteins from three independently generated datasets (*SI Appendix*, Fig. S7) (64–66). As shown in Fig. 4, 122 of the 298 (41%) HTT-interacting proteins were also found in a database of PARylated proteins. Fisher's test returned a significance of 2.1×10^{-81} . Thus, a significant proportion of HTT-interacting proteins are reported to be modified by PAR (Dataset S2).

We next asked whether HTT exists in complex with PARylated proteins in cells. Since fibroblasts do not produce sufficient protein for immunoprecipitation analyses, we turned to RPE1 cells, which are hTERT-immortalized and therefore retain DNA repair pathway function. As expected, PAR levels were increased in RPE1 cells treated with hydrogen peroxide (Fig. 4*B*). Furthermore, PARylated proteins were enriched in HTT immunoprecipitates (HTT IPs) upon oxidative stress as shown by western blot analysis. This is consistent with the high degree of overlap between datasets of HTT-interacting proteins and PARylated proteins.

To further investigate the relationship between HTT and PAR in human cells, we examined the subcellular localization of endogenous HTT upon manipulation of PAR production and degradation. While



Fig. 4. HTT interacts with PARylated proteins. (*A*) Degree of overlap between HTT interacting proteins and a list of PARylated proteins compiled from three independent studies, with Fisher's exact test for statistical significance. (*B*) RPE1 cells were treated with 400 μ M H₂O₂ in HBSS for 10 min and proteins cross-linked with 1% paraformaldehyde prior to lysis. HTT was immunoprecipitated with EPR5526 and associated proteins separated by SDS-PAGE and immunoblotted with the indicated antibodies. PARylated proteins of various sizes in the whole cell lysate (input) and anti-HTT IP were detected with pan ADP-ribose detection reagent (MABE1016) followed by HRP-conjugated anti-rabbit secondary antibody. Rabbit IgG signal from the anti-HTT immunoprecipitating antibody (EPR5526) was visible upon incubation with secondary anti-rabbit negresentative of four experiments. (*C*) RPE1 cells were treated with either 10 μ M PDD00017273 PARG inhibitor (*Top*) or 1 μ M talazoparib PARP1/2 inhibitor (*Bottom*) for 40 min prior to methanol fixation and immunofluorescence against HTT phosphorylated at residues S13 and S16 within the N17 domain (HTT phospho-N17, yellow), and PARP1 (PARP1, magenta), followed by counterstaining with Hoechst (DNA, cyan). Image representative of all mitotic cells observed (n > 10 cells from two independent experiments). (Scale bar: 10 μ m.)

endogenous HTT localization in interphase cells was similar across conditions, we found that in mitotic cells treated with PARG inhibitor, HTT phosphorylated at residues S13 and S16 localized strongly to condensed mitotic chromosomes (*SI Appendix*, Fig. S8 and Movie S1). This staining pattern is strikingly similar to that of PAR itself during mitosis (67, 68), and provides further evidence that HTT binds PAR in cells. In contrast, upon inhibition of PARP1/2 activity, HTT localized primarily to the mitotic spindle poles (Fig. 4*C*), as we and others have seen previously in untreated cells (69, 70). Thus, detection of HTT–PAR complexes by immunoprecipitation in cell lysates, and by immunofluorescence in intact cells, suggests that normal HTT may play a role in PAR biology, and raises the possibility that HTT may directly bind PAR.

HTT Contains a PBM and Directly Binds PAR. We examined the HTT sequence for potential PBMs according to consensus sequences derived by others (64, 71) and found five putative PBMs (Fig. 5*A*). While HTT putative PBMs did not match the consensus with 100% similarity, their consensus fitting was comparable to those from previously validated PAR-binding proteins including DNA-dependent protein kinase catalytic subunit (71) and 60 kD SS-A/Ro ribonucleoprotein (72).

To test these putative motifs, we performed PAR overlay assays with peptides representing the putative PBMs (Fig. 5B). PBM-3 displayed strong PAR-binding activity, which was ablated by mutation of the critical arginine residues. Multiple sequence alignment of human HTT and several orthologous species revealed a high degree of evolutionary conservation of PBM-3 and the surrounding sequences (SI Appendix, Fig. S9). Mapping of this sequence on the cryogenic electron microscopy structure of HTT (73) shows that PBM-3 is solvent accessible (Fig. 5*C*), situated on the bridge domain at the interface of the N-huntingtin, eukaryotic translation elongation factor 3, regulatory A subunit of protein phosphatase 2A, mechanistic target of rapamycin (HEAT) domain with a surface area of $\sim 890 \text{ Å}^2$. The motif spans a connecting loop region in this HEAT repeat in addition to a small section of each of the two flanking α -helices. The positively charged K1790, R1795, and R1796 residues are surface exposed in this model of the structure, indicating how they might interact with negatively charged PAR molecules.

We then asked whether the HTT PBM-3 bears resemblance to the PBM from the structurally similar (74) and functionally related (12) protein, ATM. Similar to ATM, the PBM of HTT is a surface-exposed helix-turn-helix within a HEAT repeat (*SI Appendix*, Fig. S10). These results suggest that HTT PBM-3

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ACCESSION	PROTEIN	RESIDUES	b	Х	Х	h	b	b	h	h			14 - 4
P78527	DNA-PK	2741-2748	L	Μ	Υ	А	R	Κ	G	V	F DIVI- I	Encontroll.	
P55957	BID	153-160	L	L	L	А	K	Κ	V	А			
O43684	BUB3	153-160	G	Т	А	G	R	R	V	L	PBM-2		- management
O76075	CAD	152-159	K	S	G	Y	L	R	Y	S			
P10155	SS-A/Ro	146-152	R	-	А	L	R	Κ	A		PBM-3		
Q00653	NFkB p100	179-185	K	-	Е	L	Κ	Κ	V	М			
			_	_							PBM-3 RRAA		1
P42858	HTT PBM-1	174-180	K	-	Е	T	K	Κ	Ν	G			
	HTT PBM-2	1530-1537		Α	S	G	R	Κ	A	Ŵ	PBM-4		
	HTT PBM-3	1790-1798	K	G	Μ	F	R	R	T	sт			
	HTT PBM-x	1892-1899	R	Е	I.	V	R	R	G	А		PAR	Ponceau
	HTT PBM-4	2544-2550	R	-	F	G	R	K	L	s			



Fig. 5. HTT has a PBM. (*A*) Known PBMs compared to putative PBMs in HTT. PBM-X is not solvent-accessible and was not analyzed further. b: basic, h: hydrophobic, x: any amino acid. Critical basic amino acids depicted in black boxes. (*B*) Peptides were slot-blotted onto nitrocellulose and then overlaid with 0.2 μM PAR polymer. After washing, anti-PAR western was performed with pan ADP-ribose detection reagent MABE1016. (*C, Left*) High-resolution cryoEM model of HTT-HAP40 complex (PDB–6X9O) shown in surface representation with HTT in gray, HAP40 in pink, and the PBM in orange. (*C, Right*) PBM shown in stick representation in orange. Positively charged K1790, R1795, and R1796 residues are surface exposed.

is a bona fide PBM, and that at least some of the interactions detected by immunoprecipitation and mass spectrometry (Fig. 4 and Dataset S1) could be direct.

To determine whether HTT directly interacts with PAR, we tested purified full-length HTT protein and PAR linear polymer in a fluorescence polarization assay. As shown in Fig. 6*A*, HTT binds long (26-mer) PAR chains, in preference to shorter (11-mer) substrates. This is similar to the substrate length specificity of other PAR-binding proteins (75). It should be noted that while these results support a direct interaction, the technical constraints of the experiment preclude saturation of the binding curve and therefore determination of binding kinetics and specificity. At 10 μ M protein, HTT-HAP40 Q23 exhibited lower binding than

the positive control PAR-binding protein HUWE1, but higher binding than the negative control non-PAR-binding protein USP5 (*SI Appendix*, Fig. S11). We next tested the PAR binding ability of different HTT subdomain constructs (76), and found that a construct comprising the N-HEAT and Bridge domains, which contains PBM-3, binds 26-mer PAR, while the C-HEAT domain (in complex with HAP40) does not (Fig. 6*B*). We did not detect a significant difference in PAR binding between wild-type and mutant HTT, nor upon deletion of the exon 1 domain in this assay (Fig. 6*B*). Thus, a difference in PAR binding does not explain the differential effect that wild-type and mutant HTT have on PARP1 activity in vitro (Fig. 3*C*). These results suggest that HTT can directly bind linear PAR chains of at least



Fig. 6. HTT directly binds PAR. (A) Fluorescence polarization assays using FAM-labeled 11-mer or 26-mer PAR and purified HTT-HAP40 Q23. Results for two experiments are shown. Error bars = SD. (*B*) Fluorescence polarization assays using FAM-labeled 26-mer PAR and 10 μ M of the indicated subdomains of HTT. Reactions were carried out with 3 to 4 intra-assay replicates. Results for three experiments were analyzed by one-way ANOVA and corrected for multiple comparisons by the Kruskal–Wallis test (****P* < 0.0005, ****P* < 0.0001). (*C*) Recombinant HTT-HAP40 was added to PARP1 activity assays and reactions were deposited on mica and visualized by atomic force microscopy. 2D (*Left*) and 3D (*Right*) images are shown with corresponding color scale for PARylated PARP1 (2.0 nm).

26 ADP-ribose units in length, and that binding may be mediated through PBM-3.

In order to test whether HTT may directly bind PAR chains of various sizes and branching structures, we included purified HTT protein in in vitro PARP1 reactions and visualized the reactions by atomic force microscopy at the single molecule level. In the absence of HTT, we observed auto-PARylated PARP1 structures consistent with previous reports (77–79) (Fig. 6*C*). In contrast, the presence of HTT protein in the PARP1 reactions resulted in large rosette structures consistent with HTT protein bound to PAR chains (Fig. 6*C*). While visualization of the PARP1 reaction in the presence of HTT may give insight into the increased PARP1 autoPARylation activity we detected in the presence of wild-type HTT by western (Fig. 3*C*), differing buffer conditions and mica substrate effects preclude the elucidation of precise mechanism, which is the topic of future studies.

Taken together, these results indicate that HTT binds directly to PAR chains in vitro, which accounts for its interaction with PARylated proteins in cells. This establishes a link between HTT and PAR signaling biology. Despite equivalent PAR binding, wild-type HTT increases in vitro PARP1 activity while mutant HTT does not. This may explain the reduced PAR levels in HD patient CSF, and the subdued PAR response and reduced PARP inhibitor IC₅₀ values we observed in patient-derived cells.

Discussion

The HTT protein is a large scaffold that participates in numerous cellular processes (80). We have previously defined a role for HTT in the response to oxidative DNA damage (12), and characterized the nuclear/primary cilium localization signal (81) and nuclear export signals (82, 83) regulating its translocation from the endoplasmic reticulum, where it is tethered by the N17 domain, to the nucleus (69, 84, 85). Here, we define an evolutionarily conserved PBM and show that HTT interacts with PARylated proteins. In vitro, wild-type purified recombinant HTT increases autoPAR-ylation of purified recombinant PARP1, while mutant HTT is deficient in this capacity. This may contribute to the reduced levels of PAR we observed in the CSF from HD patients and the deficient PAR levels seen in HD iPSC-derived neurons and HD patient-derived fibroblasts in the context of elevated DNA damage.

We have previously hypothesized (34, 86) that aberrant mutant HTT function in DNA repair plays a role in the elevated levels of DNA damage seen in several HD models and tissues (11–22). The ability of wild-type but not mutant HTT to stimulate PARP1 autoPARylation activity provides one possible contributing mechanism. We identified and characterized a PAR binding motif within HTT, and found that mutant HTT PAR binding was not different from that of wild type. This indicates that mutant HTT can bind PAR but cannot stimulate its activity the way that wild-type HTT can. While the identification of PBM-3 as a structurally and evolutionarily conserved PAR binding motif strongly suggests that it plays a role in HTT PAR binding, the data presented do not exclude the possibility that additional contact points contribute to PAR binding, including possibly PBM-1, which is also found in the N-HEAT subdomain and showed weak binding in the PAR overlay assay. We show that HTT directly binds PAR through the identification of a defined PAR binding motif, fluorescence polarization, and atomic force microscopy, as well as an interactome of over 100 PARylated proteins and HTT localization to PAR-coated mitotic chromosomes upon PARG inhibition.

Dysregulation of PAR signaling has now been linked to multiple neurodegenerative diseases. A common theme among these disorders is the hyperactivation of PARP1 in response to oxidative damage. This may also be the case in HD, as evidenced by strong PARP1 expression in the neurons and glia of caudate nucleus from affected HD patients (53), and by the beneficial effect of PARP1 inhibition in the R6/2 mouse model (51, 52). In contrast, we detected lower PAR levels in HD patient CSF and reduced PARP1/2 inhibitor IC₅₀ in fibroblasts from HD patients. The decreased PAR levels we detected in the CSF from HD patients, an anomaly by comparison to similar studies in different neurodegenerative diseases (32, 48–50), nonetheless reflect dysregulated PAR signaling in HD. This may parallel the paradoxical PARP inhibitor and PARG inhibitor cancer treatment options (87), which show that tipping the balance in either direction is detrimental to cancer cells. Static measurements of hypo- and hyper-PARylation may not reflect the dynamic nature of PAR metabolism, but may instead indicate PAR signaling dysregulation in HD. While CSF PAR levels did not correlate with disease measures, the robust reduction in PAR, even in premanifest patients, provides clinically relevant evidence for a role of PAR signaling dysregulation in the early stages of disease. Although we also observed lower-than-expected PAR levels in iPSC-derived neurons, the important caveat is that CSF PAR is not intraneuronal PAR. The clinical data from CSF may only indicate that there is an abnormal PAR dynamic process, hence an impaired ability to repair DNA, as seen clinically by others as early as premanifest HD (13, 14).

It is interesting to speculate on why PAR levels in the CSF are lower in HD mutation carriers while there is no difference in HD patient-derived fibroblast intracellular levels. This could reflect differences between systemic nervous system fluids versus skin-derived cells grown in a dish, whereby cell culture conditions require a minimum PAR level for cell proliferation. Basal PAR levels were in fact lower in iPSC-derived cultured, nondividing neurons representing the more severe juvenile HD case (Q77), consistent with lower PAR levels in CSF from HD patients. The ELISA method used to measure CSF PAR levels recognizes both PARylated proteins and free PAR released by brain cells during waste clearance.

The fact that PAR levels did not correlate with biomarker levels or clinical measures of disease likely reflects the transient and fluctuating nature of PAR, meaning that it may not be useful as a biomarker of disease state. This is also true of other disease-related molecules such as inflammatory markers (88) that are linked to pathology but do not have optimal kinetics for biomonitoring. Further, where PAR may be relevant to regionally specific pathology, its level in CSF is not specific to a particular brain region. Therefore, any regional-specific changes could be diluted by more general fluctuations. We speculate that dysregulation of PAR signaling in HD may have beneficial secondary consequences, such as the reduced cancer rates seen in CAG expansion carriers (89), which could possibly be related to reduced PARP1/2 activity. This could in turn contribute to the apparent evolutionary advantage of longer CAG repeat lengths, which are not in equilibrium in human populations, but are subject to mutational bias toward expansion (90).

An interaction between HD and cancer via PARP1/2 activity may provide opportunity for the repurposing of currently available cancer drugs targeting this pathway. Future studies will look at different classes of FDA-approved PARP1 inhibitor drugs as some trap PARP1 on DNA, and some can cross the blood–brain barrier while others cannot (54, 55). Interrogation of large banks of HD patient data (91) may make it possible to determine changes in disease progression associated with administration of such drugs, while the emerging development of new PARP and PARG inhibitors may hold promise for HD and neurodegenerative diseases at large.

Materials and Methods

Antibodies and Reagents. All reagents were from MilliporeSigma unless otherwise stated. Veliparib (ABT-888) was from Selleckchem. PARG inhibitor (PDD00017273) was from AdipoGen Lifesciences. Antibodies are listed in *SIAppendix, Supplementary Methods*. Mica and specimen supports were from Ted Pella Inc.

Human CSF Samples. CSF samples were collected as part of the HD-CSF study, a prospective single-site study with standardized longitudinal collection of CSF, blood, and phenotypic data (online protocol: 10.5522/04/11828448.v1). The cohort included manifest and premanifest HD mutation carriers as well as healthy controls who were age and gender matched to the entire HD mutation carrier group (*SI Appendix*, Fig. S12). All samples were deidentified prior to use in this study. In summary, lumbar punctures were performed after 12-h fasting, in the early morning, and samples were processed within 30 min after the collection. Processed samples were aliquoted and stored at -80 °C until shipment on dry ice for PAR quantification.

Measurement of CSF Analytes. PAR quantification was performed using an inhouse ELISA as previously described (50). NfL, mutant HTT, and Hemoglobin were quantified using the Simoa[®] Neurology 4-plex B kits (Item 103345, Quanterix, Lexington MA, USA, at UCL), the SMCxPro (2B7/MW1, in-house at Evotec), and ELISA (E88-134, Bethyl Laboratories; at Evotec), respectively, as described previously (57).

Cell Culture and Treatments. TruHD cells (58) were cultured in MEM (Life Technologies #10370) with 15% fetal bovine serum (FBS; Life Technologies) and 1× GlutaMAX (Life Technologies #35050) and grown at 37 °C with 5% CO₂ and 5 to 8% O₂. STHdh cells (92) were cultured in Dulbecco's Modified Eagle Medium (DMEM) (Life Technologies #11995) with 10% FBS and grown at 33 °C with 5% CO₂. RPE1 cells (American Type Culture Collection) were cultured in 1:1 DMEM/ Nutrient Mixture F-12 (DMEM/F12; Life Technologies #11330) with 10% FBS and 0.01% hygromycin and grown at 37 °C with 5% CO₂ and 5 to 8% O₂. Induction of DNA damage with H_2O_2 or KBrO3 was done in Hank's balanced salt solution (HBSS) or phosphate-buffered saline (PBS), at the concentrations and durations indicated in figure legends.

Neuronal Differentiation of iPSCs. HD iPSCs were differentiated as previously described (21, 93). All samples were deidentified prior to use in this study.

Measurement of PAR Levels in Cells. Cells were seeded in 8-well ibiTreat μ -Slides (Ibidi) to ~95% confluence. For KBrO₃ dose-response experiments, cells were seeded in glass-bottom 96-well plates (CellVis). After the indicated treatments, cells were stained and imaged as described in *SI Appendix, Supplementary Methods*.

RADD. Cells were seeded in 8-well ibiTreat μ -Slides (Ibidi) or Mod3D chambers (94) to ~95% confluence. Cells were washed with PBS and then incubated with PBS containing 0 mM or 100 mM KBrO₃ for 30 min. RADD was performed as described (59) and *Sl Appendix, Supplementary Methods*. Cells were imaged in PBS using the 20× objective on the EVOS FLAuto 2 widefield microscope. Nuclei were identified as primary objects in CellProfiler (95) using the Hoechst staining, then pixel intensity of the RADD and PAR staining within nuclei was calculated and the mean intensity recorded for each image. Eighteen images per well were captured, representing >500 cells per experiment.

Inhibitor Dose-Response Experiments. PARP1/2 and PARG activities were measured using the method described by James et al. (60) and in *SI Appendix, Supplementary Methods.* IC₅₀ values were calculated using GraphPad Prism Version 9.4.1. Means were assessed by Brown–Forsythe and Welch ANOVA tests followed by Dunnett's T3 multiple comparisons test with individual variances computed for each comparison.

Immunofluorescence. To measure chromatin retention of PARP1, TruHD cells were grown in CellCarrier-96 Ultra Microplates (Perkin Elmer) and treated for 30 min with 100 mM KBrO₃ dissolved in PBS containing Ca²⁺ and Mg²⁺. Soluble proteins were extracted with cold 0.2% Triton X-100 in PBS containing Ca²⁺ and Mg²⁺ for 2 min on ice prior to staining as described in *SI Appendix, Supplementary Methods*.

For imaging of mitotic cells, RPE1 cells were stained as described in *SI Appendix, Supplementary Methods*.

Cell Lysis and Western Analysis. For measurement of protein levels by western, cells were lysed using radioimmunoprecipitation assay buffer (50 mM Tris-HCl pH 8.0, 150 mM NaCl, 1% NP-40, 0.25% sodium deoxycholate, 1 mM EDTA) with protease and phosphatase inhibitors (Roche) and protein levels were measured with the Qubit Protein Broad Range Assay (ThermoFisher). Westerns were performed and quantified as described in *SI Appendix, Supplementary Methods*.

Purification of HTT-Interacting Proteins. STHdh^{Q7/Q7} mouse striatal precursor cells were treated with either HBSS, HBSS containing 60 μ g/mL methyl methanesulfonate for 20 min, or HBSS containing 100 μ M H₂Q₂ for 1 h. Proteins were purified as described in *SI Appendix, Supplementary Methods*.

Mass Spectrometry and Protein Identification. Samples were trypsindigested, desalted on C18 column, and liquid chromatography-tandem mass spectrometry was performed on the Q Exactive high fidelity Max mass spectrometer (SPARC BioCentre, Toronto, Canada). For database searching, tandem mass spectra were extracted by Proteome Discoverer. MS/MS samples were analyzed using Sequest (XCorr Only) as described in SI Appendix, Supplementary Methods.

PAR Overlay Assay. Peptides were ordered from Genscript (see sequences in *SI Appendix, Supplementary Methods*) and diluted to 0.5 mg/mL in PBS. One microgram of each peptide was slot-blotted onto a nitrocellulose membrane. Membranes were fixed in 2.5% glutaraldehyde/PBS for 30 min then washed with Tris-buffered saline with 0.1% Tween 20 detergent (TBS-T) (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 0.1% Tween-20), followed by incubation with 0.2 μ M PAR polymer (Trevigen) in TBS-T for 30 min, three washes with excess TBS-T, and blocking with 5% milk in TBS-T for 15 min. PAR was detected using MABE1016 (1:1,000 in TBS-T+5% milk) and antirabbit-horseradish peroxidase (HRP)(1:50,000 in TBS-T+5% milk). Membranes were then incubated with 0.5% Ponceau in 3% acetic acid for 10 min and washed in dH₂0.

Protein Purification. HTT and HAP40 constructs used in this study have been described previously (96) and in *SI Appendix, Supplementary Methods*, and are available through Addgene.

Fluorescence Polarization. FAM-labeled 11-mer and 26-mer PAR were produced as described (97). Fluorescence polarization reactions were carried out in 20 mM 2-(4-(2-hydroxyethyl)-1-piperazinyl)-ethanesulfonic acid (HEPES) pH 7.4, 50 mM KCl, 2.5% glycerol, 1 mM tris(2-carboxyethyl)phosphine, 2 mM MgCl₂, and 0.005% Tween-20 with a final concentration of 25 nM PAR substrate and 10 μ M protein in a total reaction volume of 10 μ L. Reactions were analyzed using a Synergy H4 plate reader (Biotek) and data analyzed using GraphPad Prism (version 9.4.1).

In Vitro PARP1 Reaction and Atomic Force Microscopy. For analysis by western blot, 10 fmol recombinant PARP1 was incubated in a 15-µL reaction (5 mM NAD+, 1 × activated DNA, 50 mM Tris HCl pH 7.75, 50 mM NaCl) with the indicated amounts of recombinant HTT-HAP40 for 2 h at 30 °C. Reactions were separated by sodium dodecyl-sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and immunoblotted with MABE1016 pan-ADP-ribose detection reagent. Signal intensities were quantified using ImageJ. For analysis by AFM, 35 nM PARP1 was incubated with 5 µg/mL sonicated salmon sperm DNA (Abcam ab229278) in deposition buffer (12.5 mM HEPES pH 8, 12.5 mM KCl, 1 mM dithiothreitol) in the presence of 100 μ M NAD⁺ and 10 mM MgCl₂ with or without 11.5 nM HTT-HAP40 at 37 °C for 30 min. Samples were diluted 10-fold in deposition buffer containing 3 mM MgCl₂ and 10 μ L deposited on freshly cleaved mica for 5 min before rinsing with HPLC-grade water and drying under nitrogen stream. AFM images were captured in air on a Bruker Dimension iCon (Bruker, Santa-Barbara, CA, USA) in soft tapping mode with ScanAsyst-Air tip (Bruker). In this experiment, continuous force-distance curves were recorded at 256 × 256 pixels at line rate 0.996 Hz, and the tip was oscillated in the vertical direction with an amplitude of 100 to 300 nm and at low frequency (1 to 2 kHz). Images were created using Nanoscope Analysis version 2.0 (Bruker Corporation).

Statistical Analysis.

Human CSF analyses. Statistical analysis was performed using STATA MP Version 18. *P* values of <0.05 were considered significant. CSF PAR values were assessed for normality and subsequently log transformed. We assessed potential confounders including age, gender, blood contamination, and time in the freezer. Hemoglobin was used as an indicator of blood contamination. There was an association between CSF PAR and storage time in the freezer (*SI Appendix*, Fig. S12), therefore we included storage time as a covariate in all analyses. Group

comparisons were assessed using multiple regression adding storage time as a covariate followed by postestimation Wald tests (also known as a Wald chi-square test, used to assess the significance of the coefficients in a regression model). Bonferroni correction was used to adjust for multiple comparisons. Correlations were assessed using Pearson's partial correlation.

Experiment-based analyses. All data are represented as mean \pm SEM with three independent experiments unless otherwise stated in the figure legends. Statistical analysis was performed using GraphPad Prism Version 9.4.1 and described for each experiment in figure legends. Differences among multiple means were assessed by ANOVA followed by Tukey's post hoc test. Nonnormally distributed data were analyzed with the nonparametric test (Mann–Whitney test). Assessments with P < 0.05 were considered significant. Eta squared values were used to determine gene status effect size, with 0.01 representing a small effect, 0.06 representing a moderate effect, and 0.14 representing a large effect (98).

Data, Materials, and Software Availability. All study data are included in the article and/or supporting information.

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