ABPP-HT - high-throughput activity-based profiling of deubiquitylating enzyme inhibitors in a cellular context

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12 Abstract (142 words)

13 The potency and selectivity of a small molecule inhibitor are key parameters to assess during the early stages 14 of drug discovery. In particular, it is very informative for characterizing compounds in a relevant cellular context in order to reveal potential off-target effects and drug efficacy. Activity-based probes (ABPs) are valuable tools 15 16 for that purpose, however, obtaining cellular target engagement data in a high-throughput format has been 17 particularly challenging. Here, we describe a new methodology named ABPP-HT (high-throughput-compatible 18 activity-based protein profiling), implementing a semi-automated proteomic sample preparation workflow 19 that increases the throughput capabilities of the classical ABPP workflow approximately ten times while 20 preserving its enzyme profiling characteristics. Using a panel of deubiguitylating enzyme (DUB) inhibitors, we 21 demonstrate the feasibility of ABPP-HT to provide compound selectivity profiles of endogenous DUBs in a 22 cellular context at a fraction of time as compared to previous methodologies.

23 1 Introduction

24 Activity-based probes (ABPs) can assess enzyme activity and inhibition within a cellular environment, thereby 25 providing a considerable advantage over classical biochemical and enzyme assays. ABPs typically consist of a 26 reactive group that binds to the active site of an enzyme, mostly in a covalent fashion, a specific binding 27 group/linker to aid target binding/prevent steric hindrance, and a reporter tag for fluorescence or affinity 28 (Chen et al., 2017; Chakrabarty et al., 2019; Deng et al., 2020). ABPs with an electrophilic reactive group can 29 be applied to multiple enzymes including serine hydrolases, kinases, metalloproteases and cysteine proteases 30 (Niphakis and Cravatt, 2014). Where a nucleophilic active site does not exist, photo-affinity probes can be applied instead (Niphakis and Cravatt, 2014; Mathur et al., 2020). A very informative application of these 31 32 probes is the activity-based protein profiling (ABPP), combining labeling with ABPs, immunoaffinity 33 purification, and mass spectrometry (IAP-MS) to analyze the probe interactome (Benns et al., 2021). ABPP has been used to study the potency and selectivity of small molecule inhibitors in an unbiased manner, in a cellular 34 35 matrix (Nguyen et al., 2017; Wang et al., 2018).

Profiling the ubiquitin conjugating activity of E3 ubiquitin ligases and deubiquitylating enzymes (DUBs) is of significant interest due to post-translational protein ubiquitination regulating numerous cellular pathways.

These include protein degradation, localization or controlling function (Hershko and Ciechanover, 1992; Mukhopadhyay and Riezman, 2007). Dysregulation of ubiquitination has been linked to several pathologies including cancers and neurodegenerative diseases (Popovic et al., 2014). Consequently, various E3 ligases and DUBs are being evaluated as potential drug targets for either protein targeting chimeras (PROTACs) or inhibitors, respectively, due to their proven ability to specifically target cellular protein homeostasis (Huang and Dixit, 2016; Lai and Crews, 2017; Harrigan et al., 2018).

44 DUBs offer a mechanistic entry point for probe targeting as the majority are cysteine proteases, with a smaller 45 subsection (< 15 %) functioning as metalloproteases (Komander et al., 2009; Clague et al., 2019). The activity 46 of thiol protease DUBs can be ascertained via covalent attachment of an electrophile to their nucleophilic 47 active site within the catalytic domain. This was first achieved by replacing the C-terminal glycine 76 of 48 ubiquitin with glycyl vinyl sulfone (Borodovsky et al., 2001), and further developed towards a panel of seven 49 different Ub probes with different electrophilic warheads (Borodovsky et al., 2002). These studies have provided the framework for expanding the ubiquitin-based probe concept regarding synthesis (El Oualid et al., 50 51 2010) and chemical capture (Hewings et al., 2017), but also targeting metallo-DUBs (Hameed et al., 2019) and E3 ligases (Mulder et al., 2016; Pao et al., 2016). 52

53 The inclusion of an affinity tag such as hemagglutinin (HA), FLAG, biotin, etc. to the N-terminus of a ubiquitin-54 based probe allows for DUB enrichment by immunoprecipitation (IP). Subsequent analysis by liquid 55 chromatography tandem mass spectrometry (LC-MS/MS) can identify and quantify cellular active DUBs bound 56 to the probe. This method can be used in conjunction with a DUB inhibitor to identify inhibitor potency and 57 cross-reactivity. Any DUBs that react with the inhibitor will not bind to the probe as efficiently and will be 58 reduced in the immunoprecipitated sample when compared to a control. This method was successfully applied 59 to demonstrate the selectivity of a USP7 inhibitor, with a 2-bromoethylamine warhead probe (HA-Ub-Br2). In this case, a panel of 22 DUBs were quantified (Turnbull et al., 2017). The ABPP assay was used also to assess 60 61 cellular target engagement and DUB selectivity in crude cell extracts for small molecule inhibitors against USP9X (Clancy et al., 2020), and USP28 (Ruiz et al., 2020). 62

Without fractionation the number of DUBs immunoprecipitated and quantified by MS with a propargylamine
(PA) warhead is around 30-40 (Altun et al., 2011; Ekkebus et al., 2013). To improve this methodology, we have
recently combined this probe (Ub-PA) with sample fractionation and 74 DUBs in MCF-7 breast cancer cells
were quantified. For comparison, the transcriptomics analysis of the same cells identified a very similar number
of (78) DUB mRNAs (Pinto-Fernández et al., 2019).

One of the limitations of the ABPP assay is the relatively low throughput due to the complexity of the sample preparation in proteomic applications. Here, we develop methodology to apply activity-based protein profiling in conjunction with enzyme inhibitors in a high-throughput manner, allowing for rapid screening of the concentration dependence and selectivity of multiple inhibitors simultaneously. Although this workflow can be implemented for any ABP containing an affinity tag motif, we used ubiquitin based ABPs to screen cysteine protease deubiquitylating enzymes (DUBs) and a panel of small molecule inhibitors as a methodological proof of concept.

75 2 Materials and Methods

76 Table 1: List of reagents

Reagent	Source	Identifier
ANTIBODIES		

HA (12CA5)	Roche	11666606001
USP7 (mouse)	Sigma-Aldrich	May-46
USP7 (human)	ENZO	BML-PW0540-0100
GAPDH	Invitrogen	MA5-15738
USP30	Abcam	ab235299
CHEMICALS, KITS, ENZYMES, AND OTHERS		
Sucrose	Alfa Aesar	A15583
Acid washed glass beads	Sigma-Aldrich	G4649
Sodium Chloride	Sigma-Aldrich	S5886
Glycine	Sigma-Aldrich	G7126
Tween 20	Sigma-Aldrich	P1379
Dulbecco's modified eagle medium - high glucose	Gibco	11995040
Eagles minimum essential medium	Sigma-Aldrich	M2279
Ham's F12 Nutrient Mixture	Sigma-Aldrich	N4888
Non-essential amino acids	Sigma-Aldrich	M7145
Glutamax	Gibco	35050061
Fetal Bovine Serum	Gibco	10500064
Ethanol	Merck Life Science	32221
Tris Base (Trizma)	Sigma-Aldrich	T1503
Magnesium chloride	Sigma-Aldrich	M2670
EDTA	Fischer bioreagents	6381-92-6
TFA	Sigma-Aldrich	74564-10ML-F
Urea	Sigma-Aldrich	U1250
NP-40	Sigma-Aldrich	13021
ТЕАВ	Sigma-Aldrich	T7408
PBS	Sigma-Aldrich	D8537
SDS	Sigma-Aldrich	71725
Phosphoric Acid	Fluka	79620
Formic Acid	Sigma-Aldrich	33015
Methanol	Merck Life Science	32213
Iodoacetamide	Sigma-Aldrich	11149
Dithiothreitol	Sigma-Aldrich	D9779
Acetonitrile	Millipore	1000302500
Dimethyl Sulfoxide	Sigma-Aldrich	34869
Water for Chromatography	Millipore	1153332500
BCA Protein Assay Kit	Thermo Scientific	23227
Trypsin	Worthington	LS003740 TPCK-treated
PA-W cartridges	Agilent	G5496-60000
Evotips	Evosep	EV-2001

S-Trap Plate	Protifi LCC	C02-96well
96-well microplate, U-bottom	Greiner Bio-One Ltd	650201
EXPERIMENTAL MODELS: CELL LINES		
MCF-7	ATCC	HTB-22
SH-SY5Y	ATCC	CRL2266
INSTRUMENTS		
Nanodrop	ND-1000 spectrophotometer	1000 3.8.1
SOFTWARE		
Maxquant	MPI, www.maxquant.org	Versions 1.6.10.43 & 1.6.14
Sample Prep Workbench	Agilent	Version 3.0.0
Vworks	Agilent	Version
Graph pad Prism	Prism 8	Version 8.4.3 (686)
Excel	Microsoft Office 365	

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78 Cell culture and lysis

79 MCF-7 cells were cultured in Dulbecco's Modified Eagle's Medium (DMEM) with high glucose and 80 supplemented with 10 % (v/v) Fetal Bovine Serum. SH-SY5Y cells were cultured in Eagle's Minimum Essential

81 Medium and Ham's F12 Nutrient Mix (1:1), supplemented with 15 % (v/v) Fetal Bovine Serum, 1 % (v/v) non-

essential amino acids and 2 mM Glutamax. Cells were maintained at 37 °C, 5 % CO_2 .

83 For cell collection and lysis, cells were washed with phosphate-buffered saline (PBS), scraped in fresh PBS and

84 collected at 300 xg. Cells were resuspended in lysis buffer (50mM Tris Base, 5 mM MgCl₂ · 6 H₂O, 0.5 mM EDTA,

85 250 mM Sucrose, 1 mM Dithiothreitol (DTT), pH 7.5) and vortexed with an equal volume of acid washed glass

86 beads 10 times for 30 seconds, with 2 minute breaks on ice. MCF-7 cells were clarified by 14,000 xg

87 centrifugation at 4 °C for 25 minutes. SH-SY5Y cells were clarified at 600 x G at 4 °C for 10 minutes to retain

88 USP30 bound to mitochondria. Protein concentrations were determined by BCA protein assay.

89 DUB small molecule inhibitors used in this study

USP7 inhibitors FT671 and FT827 (Ioannidis et al., 2016; Turnbull et al., 2017) were a kind gift from Stephanos
 Ioannidis. USP30 inhibitor 39 (Kluge et al., 2018), and USP30 inhibitor 3-b (patent WO2020072964) were kindly

92 provided by Jeff Schkeryantz and Lixin Qiao (Evotec/Bristol-Meyers-Squibb). Inhibitor structures are shown in

93 Figure S3. PR619 was purchased from Calbiochem (Cat. No. 662141), N-Ethylmaleimide from Sigma-Aldrich

94 (Cat. No. E3876), USP7 inhibitor P22077 from Calbiochem (Cat. No. 662142), and USP7 inhibitor HBX41108

95 from TOCRIS (Cat. No. 4285).

96 Tissue collection and lysis

97 Tissue was harvested from mice culled by exsanguination under terminal anaesthetic (isoflurane >4% in 95%O2
98 5%CO2); depth of anaesthesia was monitored by respiration rate and withdrawal reflexes. Mice were perfused

99 with PBS and tissue frozen at -80°C. Mouse brain was homogenized in lysis buffer used for the lysis of cultured

- 100 cells, using a dounce homogenizer. Once the tissue reached a homogenous consistency the glass bead lysis
- 101 protocol was carried out as outlined for cultured cells. Lysates were clarified at 600 xg at 4 °C for 10 minutes.

102 Western blotting

- 103 Samples were boiled in Laemmli sample buffer and separated on a Tris-glycine SDS page (4-15 % acrylamide
- 104 gradient) gel. Samples were then transferred to a PVDF membrane and blocked for 1 h in 4 % milk TBST.
- 105 Primary antibodies were incubated overnight at 4 °C and secondary antibodies were incubated for 1 hour at
- 106 room temperature. Imaging was carried out on a LI-COR odyssey detection system.

107 Probe synthesis

- HA-Ub-PA was synthesized as previously described (Borodovsky et al., 2002; Pinto-Fernández et al., 2019).
 Ubiquitin was expressed in *E.coli* (Gly76del) with an N-terminal HA tag and a C-terminal intein-chitin binding
- domain (CBD). *E.coli* were suspended in 50 mM Hepes, 150 mM NaCl, 0.5 mM DTT (buffer used throughout
- 111 synthesis) and sonicated 10 times, 30 seconds on, 30 seconds off. Purification was carried out using Chitin bead
- slurry, and HA-Ub-MesNa was formed via overnight agitated incubation with 100 mM MesNa at 37 °C whilst
- the protein was still attached to the Chitin beads. HA-Ub-PA was formed by incubation of HA-Ub-MesNa with
- 114 250 mM PA with agitation at room temperature for 20 minutes. Excess PA was removed via PD-10 column
- desalting. Complete and active probe formation was confirmed via anti-HA western blot and intact protein LC-
- 116 MS (data not shown).

117 Probe and inhibitor labelling

- 118 Lysates were diluted to 3.33 mg/ml using lysis buffer (minus the volume of the inhibitor and probe). Inhibitors
- 119 were diluted with either DMSO (or ethanol in the case of NEM) to the same volume for their concentration
- 120 dependence. 3-b, 39, FT671 and NEM were incubated with cell/tissue lysates for 1 hour at 37 °C. Probe was
- incubated with lysate at a ratio of 1:200 (w/w) for 45 minutes at 37 °C in all conditions, except for with USP7
- 122 (10 minute incubation) due to long probe incubations displacing bound inhibitor in this case. Reactions were
- quenched by addition of SDS to 0.4 % (w/v) and NP40 to 0.5 % (v/v) and made to 1 mg/ml protein concentration
- by addition of NP40 buffer (50 mM Tris, 0.5 % NP40 (v/v), 150 mM NaCl, 20 mM MgCl₂, pH 7.4) (freezing at
- 125 this point had no effect on the result of the subsequent IP).

126 Immunoprecipitation with Agilent Bravo AssayMAP liquid handling robot

- 127 Anti-HA (12CA5) antibody (Roche) was immobilized on Protein A (PA-W) cartridges (Agilent, G5496-60000),
- using the Immobilization App (Agilent Sample Prep Workbench v3.0.0). All steps use PBS buffer (Sigma-
- 129 Aldrich). Cartridges were primed with 100 μ L (at 300 μ L/min) and equilibrated with 50 μ L (at 10 μ L/min)
- followed by loading of 100 μ g antibody (or otherwise as stated) in a final volume of 50 μ L PBS buffer at 3
- 131 μ L/min. A cup wash with 50 μ L and an internal cartridge wash step (100 μ L at 10 μ L/min) were performed
- 132 before re-equilibrating the cartridges with 50 μ L at 10 μ L/min).
- 133 The Affinity Purification App was used for pull-downs. Briefly, Protein A cartridges with immobilized anti-HA
- antibody were primed (100 μL at 300 $\mu L/min$) and equilibrated (50 μL at 10 uL/min) with NP-40 buffer, which
- 135 was also used for all following steps. The sample was loaded at a flow-rate of 1 μ L/min. After sample loading
- 136 the cup was washed (50 μ L) and an internal cartridge wash step (100 uL at 10 μ L/min) performed to remove
- 137 unbound lysate. Peptides were eluted using 50 μL at 5 $\mu L/min$ 6M Urea or 0.15% TFA or 5 % SDS.

138 Mass spectrometry sample preparation and analysis

- 139 Urea and TFA eluates were diluted/neutralized with 180 μL 100 mM TEAB. Samples were digested in solution
- 140 with 1 μg Trypsin (Worthington, LS003740 TPCK-treated Trypsin) over night at 37 C. Digestion was stopped by
- 141 acidification to final concentration of 1% formic acid.

- 142 SDS eluates were prepared following an S-Trap 96-well plate (Protifi LLC, C02-96well) protocol. Eluates were
- acidified with ~12% phosphoric acid (10:1 v/v) and loaded into the S-trap containing 350 μ L 90% MeOH in 100
- 144 mM TEAB and spun at 1500x g for 1 min. This step was repeated three times. Then samples were resuspended
- in 100 μ L 100 mM TEAB with 1 μ g Trypsin (Worthington) and digested over night at 37 C. Samples were eluted
- from the S-traps in three consecutive steps, each for 1 min at 1500x g, first with 50 μ L 50 mM TEAB, then 50
- 147 $\,$ μL 0.1% TFA and finally 50 μL 50% ACN /0.1% TFA. The combined eluates were dried down in a vacuum
- 148 centrifuge and resuspended in 2% ACN / 0.1% TFA for LC-MS.

149 LC-MS data acquisition

- Samples were either run on a LC-MS setup comprised of an Evosep One coupled to a Bruker timsTOF Pro or a
 Dionex Ultimate3000 coupled to a Thermo Q Exactive Classic.
- 152 Evotips (Evosep) were prepared and loaded with peptides as described by the manufacturer. Briefly, Evotips
- 153 were activated by soaking in isopropanol and primed with 20 μL buffer B (ACN, 0.1% FA) by centrifugation for
- 154 1 min at 700g. Tips were soaked in isopropanol a second time and equilibrated with 20 μL buffer A (water,
- 155 0.1% FA) by centrifugation. 20 μL buffer A were loaded onto the tips and the samples were added. Tips were
- 156 spun and then washed with 20 μl buffer A followed by overlaying the C18 material in the tips with 100 μL
- 157 buffer A and a short 20 s spin.
- 158 Peptides were separated on an 8 cm analytical C18 column (PepSep, EV-1109, 3μm beads, 100 μm ID) using
- the pre-set 100 samples per day gradient on the Evosep One. MS data was acquired in PASEF mode (oTOF
- 160 control 6.2.105 / HyStar 5.1.8.1) in a mass range of 100 -1700 m/z with 4 PASEF frames (3 cycles overlap). The
- ion mobility window was set from 1/k0 0.85 Vs/cm² to 1.3 Vs/cm², ramp time 100 ms with locked duty cycle.
- 162 On the Orbitrap setup comprised of a Dionex Ultimate 3000 nano LC with Thermo Q Exactive Classic peptides
- 163 were separated on a 50-cm EasySpray column (Thermo Fisher, ES803, 2 μm beads, 75-μm ID) with a 60 minute
- 164 gradient of 2 to 35% acetonitrile in 0.1% formic acid and 5% DMSO at a flow rate of 250 nL/min.
- 165 MS1 spectra were acquired with a resolution of 70,000 and AGC target of 3e6 ions for a maximum injection
- time of 100 ms. The Top15 most abundant peaks were fragmented after isolation with a mass window of 1.6
- 167 Th at a resolution of 17,500 with a maximum injection time of 128 ms. Normalized collision energy was 28%
- 168 (HCD).
- 169 Data for experiment in Figure S1D was generated as follow: Samples of the lysate titration on cartridge have
- been run on a timsTOF Pro (OtofControl 6.0.115 / HyStar 5.0.37.1) coupled to a Dionex Ultimate 3000 on a 15
- 171 cm IonOpticks Aurora series column (1.6 μm beads, 75 μm ID, Ionopticks AUR2-15075C18A) at a flow rate of
- 400 nL/min. The gradient started for 3 min at 2% B increasing linearly in 17 min to 30% B followed by ramping
- up to 95% for 1 min and re-equilibration to 2% B. Data has been acquired in PASEF mode as described above.
- 174 Data in Figure 5 with compounds FT671, FT827, HBX41108, P22077, 3-b, 39, and PR619 has been acquired on
- a 100 samples per day gradient on a 8 cm Pepsep column (1.5 μm beads, 150 μm ID, PepSep, PSC-8-150-15-
- 176 UHP-nC) with PASEF data acquisition as described above.

177 Maxquant analysis

- 178 Orbitrap raw data was searched in Maxquant 1.6.10.43, timsTOF data was searched in Maxquant 1.6.14. MCF-
- 179 7 and SHSY5Y cell samples against a reviewed Homo sapiens Uniprot database (retrieved 31-Dec 2018), mouse
- 180 brain against a reviewed Mus musculus Uniprot database (retrieved 17-Oct 2020).

- 181 Maxquant default settings have been used with oxidation of methionine residues and acetylation of the
- 182 protein N-termini set as variable modifications and carbamidomethylation of cysteine residues as fixed
- 183 modification. The match between runs feature was used for all analyses.
- 184 Raw data and Maxquant search results have been deposited to PRIDE with the identifier PXD023036.

185 Data analysis

186 Graphs were generated and fitted using Graphpad Prism version 8.4.3 (686). All intensities from mass

- spectrometry experiments are LFQ (label-free quantitative) intensities unless otherwise stated. DUBs were filtered and removed based off presence in a no probe control sample, missing values in the probe control
- 189 sample, or intensity values that were at the bottom limit of the MS dynamic range.

190 3 Results

191 ABBP-HT workflow optimization

Each stage of the high-throughput IP methodology outlined in **Figure 1**, was optimized for maximal DUB identification coverage with minimal material to reduce experimental cost/time. Different starting material type and concentration, probe labeling, immunoaffinity purification, elution, and sample preparation conditions were tested and optimized.

First, we decided to identify the ideal ratio of probe to protein in the lysate by incubating increasing concentration of the probe with a fixed amount of lysate. Note: this step should be carried out every time after changing the batch of ABP. Our results (shown in **Figure S1A**) suggested that we should use at least 0.25 µg of HA-Ub-PA probe to label efficiently 50 µg protein extracts from two different cell lines, MCF-7 and SH-SY5Y.

In parallel, we also determined the best antibody concentration for immunoprecipitation. Anti-HA antibody was loaded onto Protein A cartridges as a set volume of 50 μ l at a flow rate of 3 μ g/uL. The column was then washed with 100 μ L of PBS buffer. From this, the loading and washing flow through fractions were collected, and unbound antibody was detected by 280 nm Nanodrop measurements. Whilst residual amounts of protein were detected at low concentrations, a significant amount of protein was present when 90 μ g was loaded, suggesting the column saturates between 80-90 μ g of antibody (**Figure S1B**). From this, we concluded that above 80-90 μ g of antibody saturates all column binding sites.

Using cartridges primed with 80 μ g of HA antibody, a concentration dependence was also carried out using varying amounts of HA-Ub-PA probe which was diluted to a set volume of 25 μ L and loaded at a flowrate of 3 μ L/min. The presence of probe in the loading and washing flow-through fractions was detected by anti-HA immunoblotting. Unbound probe was detected both in loading and washing flow-through fractions when \geq 5 μ g of probe was flowed through the column (**Figure S1C**). Therefore, no more than 2-5 μ g of probe should be

- 212 loaded to avoid column saturation and material waste.
- 213 In order to identify the optimal amount of labelled lysate we performed an LC-MS/MS analysis after performing
- 214 IAP-MS with different amounts of labelled lysate, using the parameters described above. The results (Figure
- 215 **2A**) showed that 250 μg of labelled lysate gives us the maximum of DUB identifications (IDs). Analysis of the
- flow through by immunoblotting of USP7 confirmed these results (Figure 2B, Figure S1D, and Figure S2).
- 217 The next step was to optimize the elution of the immunoprecipitated material from the column, the digestion
- 218 method, and the LC-MS/MS instrumentation. Various elution/digestion methods were trialed. Initially 6 M
- 219 Urea or 0.15 % TFA were used to elute the proteins for digestion and MS analysis. An on-column trypsin

digestion of the protein was also carried out in the presence of either HEPES buffer or 0.15 % TFA. Samples were then run on a Q Exactive Orbitrap mass spectrometer and quantified using the search software Maxquant. From this, the most efficient elution was 0.15 % TFA in combination with in-solution trypsin digestion. Different elution methods followed by in-solution digestions were then trialed on an Evosep (liquid chromatography; LC) and timsTOF Pro (mass spectrometry; MS). Comparison of 5 % SDS, 0.15 % TFA and 6M urea using this instrumentation demonstrated that 0.15% TFA is again the most efficient elution for the identification and quantification of the highest number of DUBs (**Figure 2C-E**).

227 The combined use of short gradients on an Evosep LC and the fast scan speeds using Parallel Accumulation 228 Serial Fragmentation (PASEF) data acquisition on a timsTOF Pro allowed for increased sample throughput. 229 With the preset gradients on the Evosep throughput ranges from 30-300 samples per day (SPD), compared to 230 the 6-12 SPD of common nanoflow LC setups or 9 SPD on our in-house 1 h gradient. In this experimentation 231 100 SPD is used as standard, with no increase in DUBs identified occurring from a 60 SPD method (data not 232 shown). Additionally, comparison of the Evosep One with a nanoflow LC coupled to the timsTOF Pro resulted 233 in no marked difference in the number of identified DUBs (23 vs 22 respectively) on comparable gradient 234 lengths. There is ~ 20% reduction in the number of DUBs identified with the TFA elution using the 235 Evosep/timsTOF compared to an Orbitrap MS (Figure 2D and Figure 2E) at highly reduced instrument time 236 (single run ~15 min vs 160 min). Both instruments lead to the identification of a similar panel of DUBs, with 237 some DUBs unique to each. The choice between the two instruments should balance the required DUBome 238 coverage vs throughput, and whether detection of the desired DUB is feasible with the chosen methodology.

239 Finally, since one of the applications of this methodology is DUB inhibitor characterization there is an aspect 240 of the ABP assay that should considered before testing a given inhibitor: Probe incubation time. Ub-based 241 activity-based probes, especially with the highly reactive PA (propargylamide) warhead can displace both, covalent and non-covalent, DUB inhibitors over time. As shown in Figures 2F and 2G for two different USP30 242 243 inhibitors, the reversible covalent 3-b and the reversible non-covalent 39. Increasing the incubation time 244 displaced the inhibitors, especially 3-b, from the DUB, giving the impression that the inhibitor is less potent. 245 Therefore, for reversible inhibitors we suggest minimizing incubation times with the probe. Of course, this has 246 an impact on the number of DUBs identified when performing the ABPP assay. We compared two labelling 247 times using our ABPP-HT workflow and as expected, the intensities of some DUBs are clearly reduced when the lysate is incubated with probe for a shorter period of time (10 minutes versus 45; Figure 2 H). These 248 249 optimization steps were summarized in Figure 2I.

250 Guide to DUB picking: abundance changes with methodology and starting material source.

251 These conditions were then applied to characterize the active DUBome in two different cell lines, MCF-7 and 252 SH-SY5Y, and brain tissue material from mice. We also included the data using the two different LC-MS/MS instrumentations: Nanoflow liquid chromatography coupled to an Orbitrap MS (OT on the figures) and 253 254 microflow (Evosep) liquid chromatography and ion mobility-mass spectrometer, timsTOF (TT on the figures). 255 We summarized the results in a heat map (Figure 3), displaying the normalized intensities of the identified 256 DUBs when using different cell lines, tissue, and instrumentation. This together with the heat maps describing 257 the different elution methods (Figures 2D and 2E) should be a good reference when studying a particular DUB 258 and its potential inhibitors. For example, there are some DUBs that are only identified in the cancer cells like 259 USP3, USP4, and OTUD7B. On the other hand, there are DUBs specific for brain tissue and cells like UCHL1.

260 Proof of concept: Broad and specific DUB inhibitor concentration dependences and cross-reactivities

261 The ABPP-HT methodology is able to identify a representative panel of DUBs that is comparable to the regular ABPP (~15-25 vs ~30-40) (Pinto-Fernández et al., 2019). With this representative panel we applied the 262 263 methodology to check for compatibility with DUB inhibitor characterization. In order to do so, we performed ABPP-HT with the highly selective USP7 inhibitor FT671 (Turnbull et al., 2017) and with the broad cysteine 264 265 modifier NEM (n-ethylmaleimide) (Pinto-Fernández et al., 2019). We performed these experiments treating lysates from MCF-7 cells and from mouse brain tissue extracts with different concentrations of the inhibitors. 266 We used the TT LC-MS/MS instrumentation as it is more suitable for testing a higher number of compounds. 267 268 First, we performed control immunoblots against USP7, to show the effects of the compounds on USP7, and 269 against the probe (anti-HA), to visualize the selectivity of the compounds. FT671 inhibits USP7 in both, cell line 270 (Figure 4A) and brain tissue (Figure 4B), however, due to the USP7 antibody recognizing USP7 with and without 271 its previously characterized ubiquitination (Fernández-Montalván et al., 2007), in the mouse tissue this effect 272 was more challenging to visualize. At the same time, the HA blot showed little reactivity of FT671 with other 273 labelled DUBs. On the other hand, NEM was also inhibiting USP7 in both cells (Figure 4C) and brain (Figure 4D), 274 however, in a non-selective way as shown by the overall decrease in HA signal at high concentrations of the 275 compound. These observations were highly comparable when processing these samples on our ABPP-HT 276 workflow. For instance, immunoblot densitometry of labelled USP7 with increasing amounts of FT671 277 correlates to a similar degree with the LC-MS/MS data (MCF-7 on Figure 4E and brain on Figure 4G). This was 278 also the case for the selectivity profile of the two compounds in both cells and brain (Figures 4F, 4H, 4I, and 279 **4J**), reflecting well both, the expected high selectivity of FT671 and the broad inhibition by NEM.

Exploiting the possibilities of the ABBP-HT methodology: multiple compound characterization in different cell lines and tissue

282 Finally, we decided to gain advantage of the ABPP-HT possibilities and applied the methodology to a number of compounds (structures in Figure S3) and concentrations simultaneously. Here, critical target engagement 283 284 information was obtained in a cellular context, in a much faster way than the current methodology. We tested 285 different concentrations of four USP7 inhibitors, FT671, FT827 (Turnbull et al., 2017), HBX41108 (Colland et 286 al., 2009), P22077 (Altun et al., 2011), two USP30 inhibitors (3-b and 39), and the two broad cysteine modifiers 287 NEM (Pinto-Fernández et al., 2019) and PR619 (Altun et al., 2011). The results are summarized on separated 288 heat maps for USP7 inhibitors (Figure 5A), USP30 (Figure 5B) and non-selective (Figure 5C), and bar graphs (Figures S4E-H). These results not only match the matching control immunoblots in Figure S4 (A-D) but also 289 290 previously reported information. For instance, P22077 was reported to be a dual USP7/USP47 inhibitor (Altun 291 et al., 2011) and the same result could be seen in our data (Figure 5A). FT671 and FT827 were reported to be 292 highly selective, and potent, USP7 inhibitors (Turnbull et al., 2017) and this still applied when using our ABPP-293 HT workflow (Figure 5A). HBX41108 selectivity data has not been reported, although our results suggested a 294 not very selective profile. The USP30 inhibitors showed a nice dose-dependent inhibition of the target and 295 good selectivity profiles, especially for the USP30 inhibitor 39 (Figure 5B). Finally, the two cysteine modifiers 296 behaved as expected (Altun et al., 2011) (Pinto-Fernández et al., 2019), inhibiting all the identified DUBs at 297 high concentrations (Figure 5C). The ability to analyze multiple concentrations also allowed for the plotting 298 and determination of the half-maximal inhibitory concentrations (IC50s) for each inhibitor in the described 299 conditions (Figure 5D-J).

300 4 Discussion

There are numerous methods to study the potency and selectivity of an enzyme inhibitor using recombinant purified proteins, and their substrates, in biochemical assays. However, these assays cannot assess the activity

303 of an inhibitor in a more relevant context such as cell lysates, intact cells or tissue. Degradation, limited

permeability, or cross-reactivity of an inhibitor in the cellular environment may lead to reduced potency and off-target effects. Consequently, it is important to be able to screen potential inhibitors within this environment. ABPP assays can provide all of these very relevant parameters, however, if we want to apply this technique to a screen of inhibitors with varying concentrations in different cell types, the throughput needs to be increased.

309 Here, we describe a new ABPP methodology, named ABPP-HT (high-throughput-compatible activity-based protein profiling), that allowed the semi-automated analysis of samples in a microplate format, addressing the 310 low throughput associated to the classic ABPP assay. The incorporation of a liquid handling robot compatible 311 312 with IAP-MS, and the Evosep/timsTOF LC-MS/MS instrumentation, were key to boost the throughput of the 313 ABPP up to ten times in a cost-effective way. The depth of this method is reduced but comparable to the 314 normal ABPP, with the detection of ~15-25 DUBs when using the ABPP-HT versus ~30-40 DUBs with the original 315 ABPP. The number of DUBs that are reactive with the probe and can be potentially detected by ABPP is higher than 70, but this requires performing a high-pH pre-fractionation of the samples prior LC-MS/MS analysis 316 317 (Pinto-Fernández et al., 2019). This drastically increases the number of samples to analyze per condition and 318 therefore the required time and cost of the assay. This comparative information has been summarized in Figure 6. The methodology of the ABPP-HT approach can be applied as a powerful initial screening tool for 319 320 multiple inhibitors at different concentrations, in various cell lines, to discard weak or highly cross-reactive 321 inhibitors quickly and robustly. From this, only potent and selective inhibitors could be taken forward for more 322 thorough characterization using the original or fractionated ABPP approaches.

As a proof of concept, we demonstrated the versatility of this methodology using general and specific DUB 323 324 inhibitors in two different cell lines and mouse brain tissue. ABPP-HT permitted the simultaneous analysis of 6 325 selective DUB inhibitors and the calculation of their respective half maximal inhibitory concentration (IC50) 326 values. The methodology has the capacity to simultaneously test a much higher number of inhibitors and 327 concentrations. We also believe that the throughput of the ABPP-HT can be increased even further. For 328 example, by implementing chemical labels, such as TMT (tandem mass tag) that would allow the combination 329 of up to 16 samples into one and therefore providing multiplexing capabilities and enhanced throughput. 330 Another area where the sensitivity and therefore DUB coverage of this type of analysis could be further 331 improved is by implementing targeted proteomics methods such as Data-Independent Acquisition (DIA) mass 332 spectrometry. For instance, DIA has been successfully applied for ubiquitomics and discussed by Vere et al. (Vere et al., 2020). 333

In conclusion, ABPP-HT (high-throughput-compatible activity-based protein profiling) was conceptualized, optimized, and validated. When tested, the approach allowed for reduced time and cost for both sample preparation and MS time, whilst still identifying and quantifying a representative panel of endogenously expressed DUBs, enabling the profiling of a number of DUB inhibitors.

338 5 Conflict of Interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

341 6 Author Contributions

H.J., R.F., B.M.K. and A.P.F. directed this study. Most experiments were devised by H.J., R.H., B.M.K. and A.P.F.

and carried out by H.J, R.H. and A.P.F. H.J., R.H., B.M.K. and A.P.F. wrote the paper. All authors commented on
the text.

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- 451
- 452 Data Availability Statement

The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE (Perez-Riverol et al., 2019) partner repository with the data set identifier PXD023036.

455 Ethical Statement for studies involving animal subjects

The breeding of mice was carried out in accordance with Animal [Scientific Procedures] Act 1986, with procedures reviewed by the University of Oxford clinical medicine animal care and ethical review body (AWERB), and conducted under project licenses PPL POC27F69A. All procedures conformed to the Directive 2010/63/EU of the European Parliament.

Tissue was harvested from mice culled by exsanguination under terminal anaesthetic (isoflurane >4% in 95%O2
 5%CO2); depth of anaesthesia was monitored by respiration rate and withdrawal reflexes. Mice were perfused
 with PBS and tissue frozen at -80°C.

463 Figure legends

464

Figure 1. Accelerated DUB inhibitor ABP IP workflow. A. Protein extraction and inhibitor treatment of either
 intact or lysed tissue/cell lines. B. HA-Ub-PA probe incubation to label uninhibited cysteine active DUBs. C.
 Anti-HA IP, traditionally with centrifugation or magnetic collection of agarose beads in a low throughput
 format. In this work the throughput is increased to a 96 well plate format using an Agilent bravo liquid handling
 platform. D. LC-MS/MS proteomic analysis of immunoprecipitated DUBs. Here we compare the depth of the
 DUBome obtained using a QE orbitrap vs. a high-throughput timsTOF.

471

472 Figure 2. Optimizing the ABPP-HT workflow. A. Number of DUBS identified by timsTOF MS with increasing 473 amounts of HA-Ub-PA-labelled MCF-7 lysate protein, after immunoprecipitation and elution with 0.15 % TFA. 474 **B.** Western blot densitometry quantification (full blot in Figure S1D) of USP7 in the immunoprecipitation loading flow-through, with increasing amounts of HA-Ub-PA-labelled MCF-7 lysate protein quantity 475 476 immunoprecipitated and eluted with 0.15 % TFA. C. Number of DUBs identified by LC-MS/MS with different IP 477 elutions: 0.15 % TFA, 6 M urea, HEPES, *= On column digestion. D. Log2 intensities of DUBS identified with 478 different elution methods by a QE orbitrap MS. 0.15 % TFA, 6 M urea, HEPES, *= On column digestion. E. Log2 479 intensities of DUBs identified with different elution methods by a timsTOF MS. 0.15 % TFA, 6 M urea, 5 % SDS. 480 F. USP30 immunoblots showing mouse brain lysate displacement of a covalent (3-b) and non-covalent (39) 481 USP30 inhibitor with increasing HA-Ub-PA (at 37 °C) incubation times. G. The densitometric quantification of Figure 2F from the intensity of the HA-Ub-PA-labelled band, normalised to the intensity of both USP30 bands 482 483 together. H. timsTOF DUB intensities of MCF-7 labelled with HA-Ub-PA for 10 minutes normalised to 45 484 minutes at 37 °C (SEM, n=3). I. Optimisation workflow for high-throughput DUB inhibitor screening using ABPP 485 LC-MS/MS.

486

Figure 3: ABPP-HT reveals cell type-specific DUB profiles. DUB intensities as determined by HA-Ub-PA activity based probe profiling (ABPP) and identified from different cell types and tissue quantified by mass
 spectrometry using either a QE orbitrap (OT) or timsTOF (TT), normalised within each dataset.

490

491 Figure 4. ABPP-HT allows fast generation of DUB inhibitor selectivity profiles in a cellular context. A-D. USP7,

492 HA, and GAPDH (loading control) immunoblots of a concentration dependence of FT671 USP7 specific inhibitor

493 and NEM (a general cysteine modifier) in mouse brain tissue and MCF-7 cell lysates. E. The densitometric

- 494 quantification (WB) of three independent experiments as in Figure 4A from the intensity of the HA-Ub-PA USP7
- 495 MCF-7 labelled band (SEM, n=3), normalised to the intensity of both USP7 bands together, compared to the

496 MCF-7 timsTOF LFQ normalised intensity (TT) of immunoprecipitated USP7 (MS data SEM n=3 (for 0.2 μ M, 497 n=2)). F. The activity of a panel of DUBS identified by timsTOF MS from MCF-7 with increasing concentration of FT671 (SEM n=3 (for 0.2 µM, n=2)). G. The densitometric quantification of Figure 4B from the intensity of 498 499 the HA-Ub-PA USP7 mouse brain labelled band, normalised to the intensity of both USP7 bands together (WB), 500 compared to the mouse brain timsTOF LFQ normalised intensity (TT) of immunoprecipitated USP7. H. The activity of a panel of DUBS identified by timsTOF MS from mouse brain with increasing concentration of FT671. 501 502 I-J. The activity of a panel of DUBS identified by timsTOF MS from mouse brain and MCF-7 lysates respectively, 503 treated with either 1 mM or 10 mM of NEM.

504

505 Figure 5. ABPP-HT reveals DUB inhibitor selectivity and specificity compatible with higher throughput. A. The 506 activity of a panel of DUBs from MCF-7 identified from timsTOF MS, in response to USP7 specific inhibitors 507 FT671 (n=3 (for 0.2 μM n=2)), FT827, HBX108 and P22077. **B.** The activity of a panel of DUBs from mouse brain 508 lysate identified from timsTOF MS, in response to USP30 specific inhibitors 3-b and 39. C. The activity of a panel 509 of DUBs in MCF-7 lysates identified by timsTOF LC-MS/MS, in response to the cysteine modifier NEM, and 510 broad spectrum DUB inhibitor PR619 (PR619 n=2). D-I. From left to right concentration dependences of USP7 511 from inhibitors FT671, FT827, HBX41108, and P22077 in MCF-7 lysates, and USP30 inhibitors for 3-b and 39 in 512 mouse brain. K. IC50 values extracted from D-I, fit to equation: Y=100/(1+X/IC50). * = normalised raw 513 intensities, not LFQ intensities.

514

520

Figure 6. ABPP workflows optimised for DUBome depth and throughput. Comparison of cost and time
between ABPP assays optimised for maximal coverage of cellular DUBs (DUBome depth – Y-axis) and higherthroughput approaches including traditional ABPP, immunoblot based ABPP and accelerated throughput ABPP
(ABPP- HT) (Throughput – X-axis). Size of the circles are plotted represent relative cost per inhibitor compound
tested.

521 Figure S1. Optimizing DUB target engagement in the ABPP-HT workflow A. HA immunoblot of probe to lysate 522 ratio optimisation, with 50 µg of either SHSY5Y lysate or MCF-7 lysate incubated with increasing amounts of 523 HA-Ub-PA at 37 °C for 45 minutes. B. Concentration dependence of unbound antibody detected with antibody 524 loading onto protein A columns. C. Concentration dependence of unbound HA-Ub-PA probe detected with HA-525 Ub-PA loading onto protein G column with 80 µg of antibody loaded. **D.** USP7 detected in IP flow-through 526 where 100 µg of HA antibody has been loaded. Values in red were used for quantitation in figure 2B. ^a protein 527 A column, ^g protein G column, *material was concentrated before loading to try to minimize loading time, the 528 reduction in material is attributable to protein lost during the concentration step, not to increased HA-Ub-PA anti-HA binding. In each lane 5 ug of protein was loaded except in the case of 10 µg and 50 µg where samples 529 530 were too dilute to load 5 µg.

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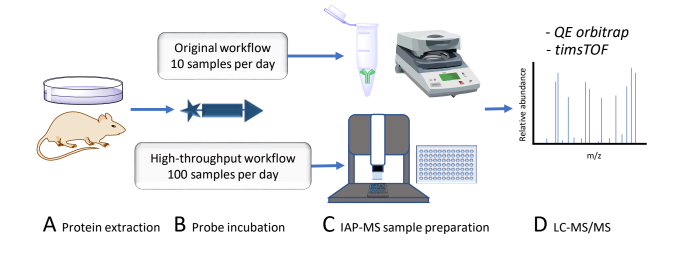
Figure S2. Efficient immunoprecipitation of USP7 with ABPP-HT USP7 immunoblot of FT671 concentration
 dependence. Before immunoprecipitation demonstrates probe labelling and inhibition, the supernatant and
 eluant bands show unlabelled and labelled bands respectively.

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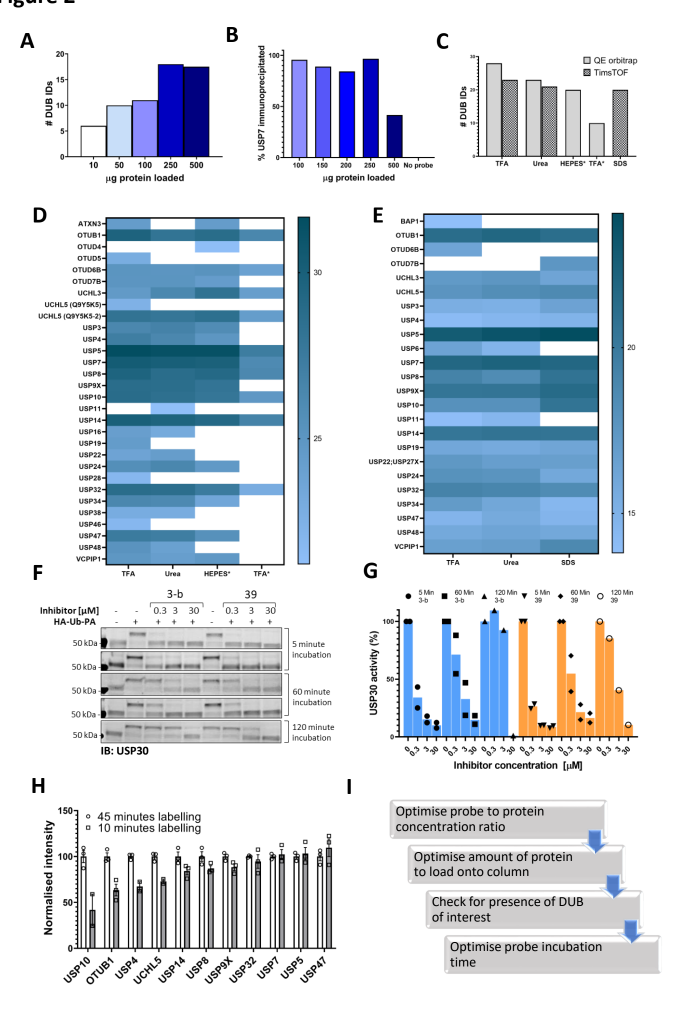
Figure S3. DUB inhibitors used in this study. USP7 inhibitors FT671 FT827, P22077, HBX 41108, USP30 inhibitor
39, USP30 inhibitor 3-b, and broad-spectrum cysteine modifiers NEM and PR619.

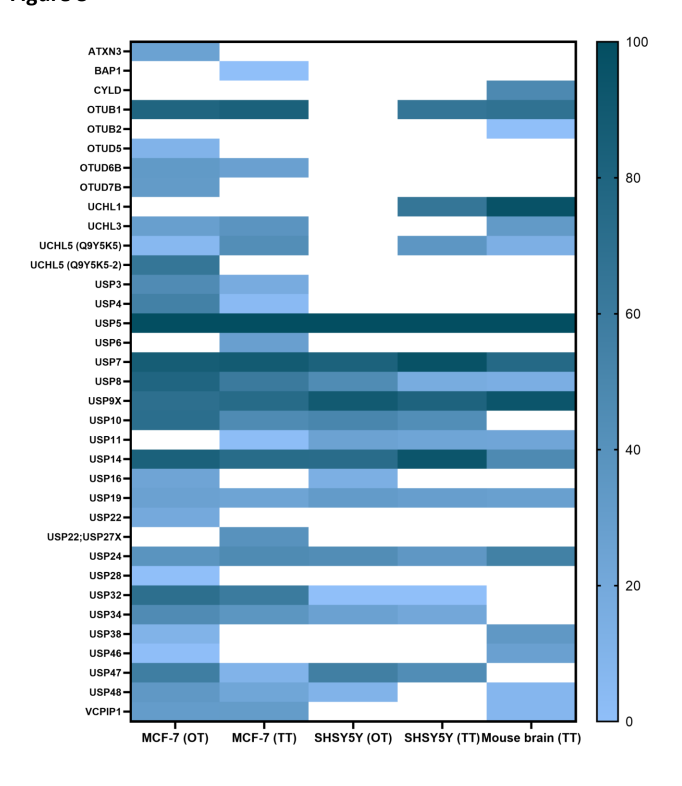
Figure S4. Inhibitor cellular target engagement assessed by ABPP. A. USP30 immunoblot of specific USP30
 inhibitors 3-b and 39, and broad DUB inhibitor NEM in mouse brain lysates. B. USP7 and anti-HA immunoblot

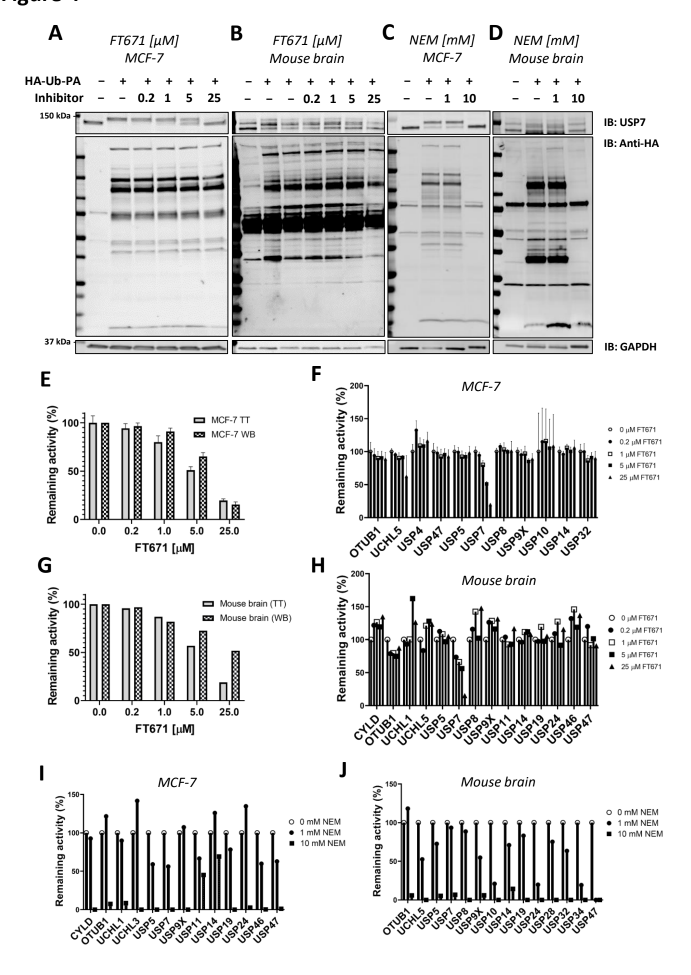
- of specific USP7 inhibitors FT829, HBX108 and P22077. C. USP7 and anti-HA immunoblot of specific USP7
- 542 inhibitor FT671. **D.** Anti-HA immunoblot with broad-spectrum DUB inhibitor PR619. **E-H.** ABPP inhibition profile
- of a panel of DUBs by FT827, HBX41108, P22077, and PR609 (one experiment, except for PR609: SEM; n=2).

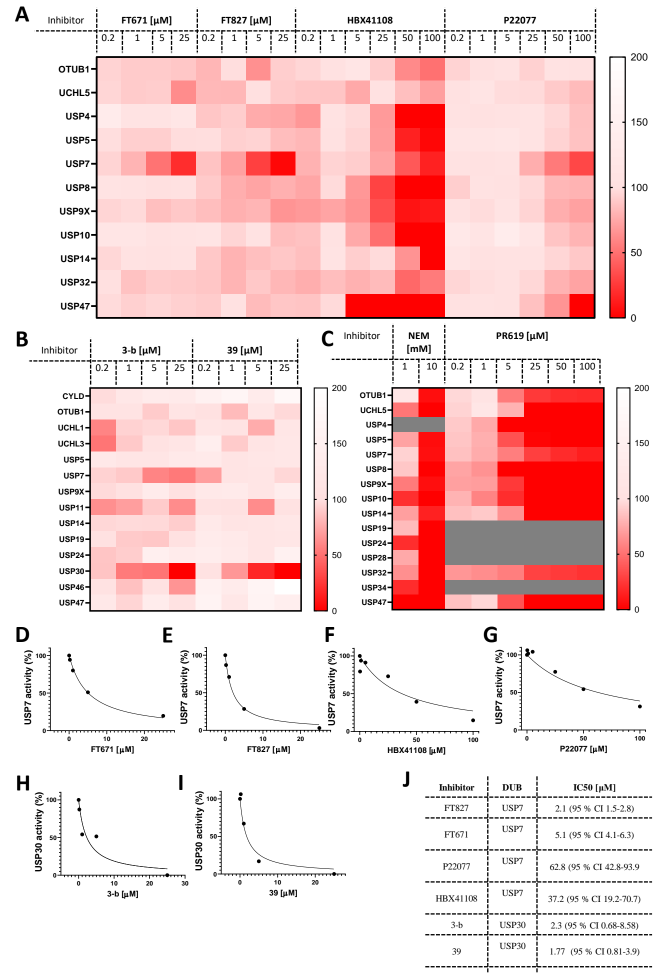


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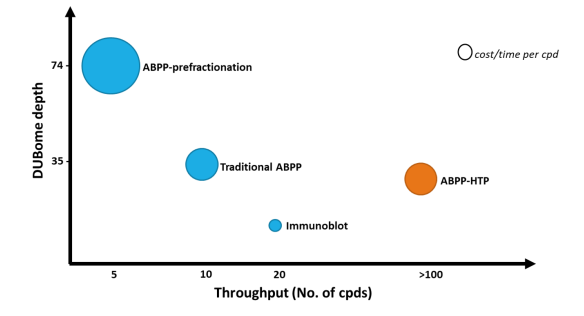


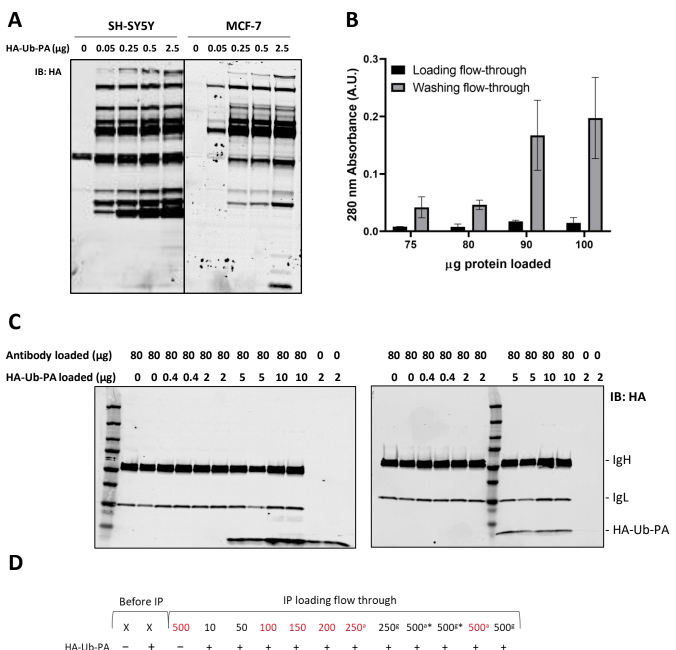


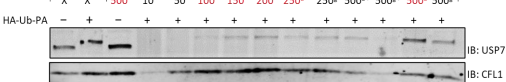




PR619 USP7 7 (95 % CI 3.6 to 13.2)







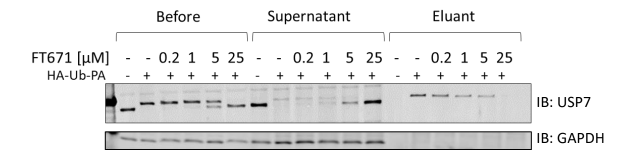
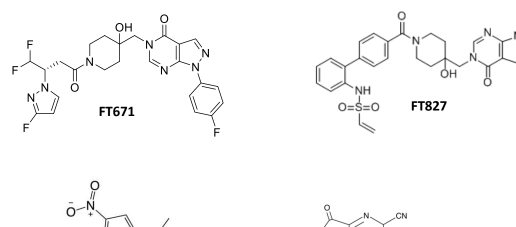
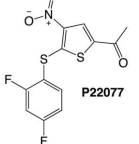
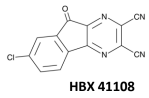
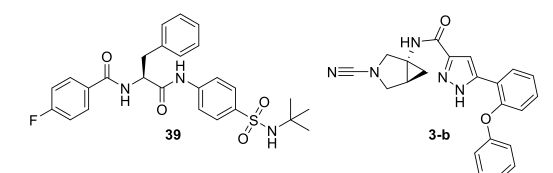


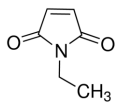
Figure S3 Figure solution for the second sec

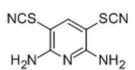












NEM

PR619

