SARS-CoV-2 interactome with Human Ghost proteome: A neglected world encompassing a wealth of biological data

Tristan Cardon¹*, Isabelle Fournier¹,²* and Michel Salzet¹,²*

¹Univ. Lille, Inserm, CHU Lille, U1192 - Protéomique Réponse Inflammatoire Spectrométrie de Masse - PRISM, F-59000 Lille, France
²Institut Universitaire de France, Paris, France

Abstract:
Conventionally, eukaryotic mRNAs were thought to be monocistronic, leading to the translation of a single protein. However, large-scale proteomics has led to a massive identification of proteins translated from mRNAs of alternative ORF (AltORFs), in addition to the predicted proteins issued from the reference ORF or from ncRNAs. These alternative proteins (AltProts) are not represented in the conventional protein databases and this “Ghost proteome” was not considered until recently. Some of these proteins are functional and there is growing evidence that they are involved in central functions in physiological and physiopathological context. Based on our experience with AltProts we have got interested in finding out their involvement in development of the SARS-CoV-2 virus, responsible for the 2020 Covid-19 outbreak. Thus, we have scrutinized the recently published data by Krogan and coworkers (2020) on the SARS-CoV-2 interactome with host cells by co-IP in the perspective of drug repurposing. The initial work has revealed the interaction between 332 human cellular RefProts with the 27 viral proteins. Re-interrogation of this data using 23 viral targets and including AltProts, followed by enrichment of the interaction networks, leads to identify 218 RefProts (in common to initial study) plus 56 AltProts involved in 93 interactions. This demonstrates the necessity to take into account the Ghost proteome for discovering new therapeutic targets and establish new therapeutic strategies. Missing the ghost proteome in the drug metabolism and pharmacokinetic (DMPK) drug development pipeline will certainly be a major limitation to the establishment of efficient therapies.

INTRODUCTION

Because proteins are the end products of gene expression, they have a major impact on cell regulation thus being main targets for the development of new drugs and therapies. Therefore, holistic approaches must be developed to grasp the proteome in its completeness and find out how it relates to the upstream genes it is issued from. Grasping the proteome can be difficult because of the broad dynamic ranges it spans on (i.e. >7 orders of magnitude from 1 copy to up to 10 million per cell) when compared to transcriptome (only 3-4 orders of magnitude) [1]. However, thanks to the last generation of liquid chromatography-mass spectrometry (LC-MS) instrumentation, >5,000 proteins can be identified in a single run experiment by large-scale bottom-up proteomics [2]. Both bottom-up and top-down proteomic approaches are very powerful; though they do show a major drawback since the protein identification is based on databank interrogation. Databanks are thus critical to large-scale proteomic approaches since only proteins referenced in the database can be identified. A large part of the proteins in databases, such as for UniprotKB/SwissProt which is the reference database in proteomics [3]; is predicted from genes according to well established rules. Thereof, only >100 codon sequences of mRNA starting with an “AUG” and presenting the favorable consensus Kozak motive are translated into a single protein accordingly to the admitted idea that eukaryotes are...
monocistronic. The single protein product expected from gene translation is designated as the reference protein (RefProt).

However, eukaryotic translation was finally demonstrated to be polycistronic as already suspected in the late 1990’s by M. Kozak [4]. Indeed, alternative translation mechanisms, such as the reinitiation or the leaky-scanning, leading to translation from alternative ORFs (AltORFs) were already described by that time; though those has remained considered as an epiphenomenon. Hence, a huge number of proteins were lacking from protein databases and have simply remained invisible to all proteomic studies, representing thereby a ghost proteome. This ghost proteome was eventually unveiled by two distinct approaches, one using ribosome profiling [5], and the second, MS-based proteomics. In ribosome profiling, many possible fixations of ribosome were described from ncRNA and untranslated region (UTR) of mRNA [6,7] highlighting the existence of non-expected protein products in mammals. From proteomic data, by using novel databases that were including protein predictions translated from AltORFs novel protein sequences were identified, filling the gap of good quality data remaining unmatched after conventional database interrogation (>10% data) [8]. These proteins, designed as alternative proteins (AltProts), are neither proteoforms, nor proteins issued from alternative splicing. Some show sequence similarities with proteins carried by other mRNA, but the others present totally new amino acid sequences. Finally, identified AltProts are found to be translated, either from mRNA including from the non-coding 5’ & 3’ UTR or a frame shift (+1 or 2 nucleotides) in the CDS of the RefProt, or from non-coding RNA (ncRNA) [9]. Overall, large-scale bottom-up [9–12] and top-down [13,14] proteomics have enable the identification of an important number of these AltProts. Very importantly, AltProts were also shown to be functional and carrying important cell functions [12,15–17]. In a way, the rediscovery of the “lost world” of protein products, will open a new page in the history of biological mechanisms. A total of ~450,000 proteins has ultimately been predicted in human and are publicly available through the OpenProt [18] database. This is about 20-fold more than yet estimated from conventional databases (20353 entries in June 2020 for reviewed RefProt). It is thus possible to gain incredible knowledge by considering AltProts in already generated data. Previously, proteomic data reuse, have enabled the discovery of the ghost proteome interactome using cross-linking MS (XL-MS) data from HeLa cells [19,20]. In this study, AltProts were found to be interacting with RefProts involved in protein translation regulation as evidenced by the participation of AltATAD2 in the RPL10/AUF1 complex [20]. Since, the study of glioma cell line (NCH82) under activation by a protein kinase A activator inducing a cellular phenotypic change, has confirmed the presence of AltProts in the signaling pathways of protein translation. AltProts were also shown interacting with cytoskeleton proteins (e.g. AltTRNAU1AP, AltMAP2 and AltEPHA5 interacting with TPM4) [10].

Based on our experience with AltProts we have got interested in finding out their involvement in development of the SARS-CoV-2 virus, responsible for the 2020 Covid-19 outbreak. Thus, we have scrutinized the recently published data by Krogan and coworkers [21] on the SARS-CoV-2 interactome with host cells by co-IP in the perspective of drug repurposing.

MATERIAL AND METHODS

1. Ghost Proteins Databases.

The study was carried out using OpenProt database (www.openprot.org) [18,22]. This database is derived from the predicted H. Sapiens alternative proteins (GRCh38.p5, Assembly: GCA_000001405.20). This database compile all proteins coming from non-coding regions of mRNA such as 5′&3′ UTR, shift in reading frame in +2 or +3, and the proteins discovered coding in ncRNA. Also to this database the RefProt from UniProtKB is added, for a total of 658263 entries. ProteomeDiscover 2.3 (PD2.3) with label free quantification node is used to analyse the RAW data from ProteomeXchange consortium via PRIDE repository dataset,
Following parameters is apply on PD2.3: trypsin as enzyme, 2 missed cleavages, methionine oxidation as variable modification and carbamidomethylation of cysteines as static modification, Precursor Mass Tolerance: 10 ppm and fragment mass tolerance: 0.6 Da. The validation was performed using Percolator with a FDR set to 0.001%. A consensus workflow was then applied for the statistical arrangement, using the high confidence protein identification and at least one unique peptide for identified proteins.

The identified proteins, are correlated with the bait of CoIP describe on the dataset and to the PRIDE project [21]. Protein identified with a fold change up to 2 between the bait expression and the control of CoIP are keeping as potential interactor. The network is drawing on Cytoscape V.3.8.0 [23], the DyNet [24] application is used to compare the network publish in NDEx (according to [21]) and our result. A color code is given for nodes: red hexagon are the viral protein (bait), blue circles are the RefProts and green circles are the AltProts, and for the edges: red are interaction not recovered in our analysis, grey are recovered in both analysis, green are specific to our analysis and with a ratio<100 when purple edges are interaction specific to our analysis with a ratio of 100. A ratio of 100 meaning that protein is not detected in the control, and the expression can be link to the expression of the viral protein. The AltProt identified have been describe based on the recovered information obtain from OpenProt database, Ensembl and RefSeq database.

Blast analysis (non-redundant sequences and RefSeq) of the AltProts sequences, identified in interaction with the SARS-CoV-2 proteins, show the presence of 27 AltProts exhibiting a homology rate greater than 80% (average of the coverage and identity percentage). These proteins, for the major part are ncRNAs emitted, are therefore not isoforms of homologous proteins because they originate from a different RNA sequence. From the total list of AltProts identified, Blast analysis reveal 16 AltProts with no significant (<80%) homologies, these 16 can have a known protein domain based on few identity with referenced protein, but experimental data is need to proof the context of action to this AltProt. In the same way, 16 other AltProts have no Blast result in human database (non-redundant sequences and RefSeq).

In the context of following and understanding the SARS-CoV-2 way of action in the host cell, and considering the bat origin of the virus, the proteins sequences of the no result blast has been interrogated to the bats database (taxid:9397). 7 of the 16 AltProts describe similarity in bats protein, with a rate between 35 and 78% of homology.

Results and discussion

We have searched for interactions between host AltProts and the viral SARS-CoV-2 proteins to study the impact of the ghost proteome in the viral interaction to host cells. The SARS-CoV-2 virus, expresses a 30 kb genome capable of coding for at least 14 ORFs, capable of producing more than 27 proteins[25]. The initial work[21] has revealed the interaction between 332 human cellular RefProts with the 27 viral proteins. Re-interrogation of these data using 23 viral targets (those not found at the membrane) and including AltProts database, leads to identify 218 RefProts (common with the initial study) plus 56 AltProts involved in 93 interactions (Figure 1) of which 17 interacted with more than one viral proteins. 59% of them originate from ncRNA, 41% from mRNA of which 39% from 3’UTR region, 34% from 5’UTR region and 26% from a CDS shift (Table.1). 26 AltProts are exclusive to the host cells stimulated by the viral protein and not found in the control. Some identified proteins and interactions are found to be different from the initial study because a different methodology was applied in the data reuse. This is a consequence of using a larger size database including both RefProts and AltProts, then forcing the utilization of Proteome Discoverer in place of Maxquant, following the recommendations of the OpenProt developers[18,26]. However, strong FDR filter is used,
unique peptide is verified for each identified protein, and a cutoff threshold sample/control of 2 is applied to define an interactor.

25 AltProts, after a Blast using human nun redundant database, present a strong homology (>80% of the average percentage of coverage and percentage of identity) to a RefProt, though they are identified with a unique peptide to the AltProt sequence. This case are not isoform because coming from another gene of the RefProt or from an ncRNA, but share a common domain with the referenced or predicted protein. Global analysis of the Biological Processes of proteins identified as homologs shows that the mainly pathway impacted the protein metabolism (Fig.2-A), in particular with signaling pathways such as protein translation and elongation (EIF2S2; EEF1A1; RPL35A; RPL4 ; RPS17; RPS18; RPL18A) and the regulation of protein synthesis by insulin (UBE2D3; HSPD1; HSPA8; PRKDC; HNRNPA1) interestingly proteins: RPL35A; RPL4; RPS17; RPS18; RPL18A; are found in the biological process of viral RNA translation, and particularly in influenza.

Interestingly, it has been described that SARS-CoV-2 proteins were impacting the phosphorylation state of the host cell proteins, like the N protein which was shown to differentially phosphorylate LARP1 and RRP9 [27]. In this way it’s not a surprised to recover some AltProt with riboprotein domain in interaction with SARS-CoV-2 proteins like IP_668819, IP_637436, IP_639311, IP_597129, IP_750273 and IP_667059. These protein are identified to be in interaction with the non-structural protein nsp8 (IP_637436, IP_750273, IP_667059), nsp12 (IP_639311) two viral protein described to be involved in the virus RNA replication [28–30]. So finding interaction with ribosomal protein and AltProt is not a surprised, as the nsp8 and nsp12 complex on the RNA when the ribosomal protein work at the translation in protein. In fact, more than 37 ribosomal protein (RPL) can be observed in interaction with nsp8, RefProt and AltProt confounding.

The major observation of the Bats homology with the no result human Blast analysis, show the importance of the AltProt studies, indeed if today an important part of the protein is knowing and identified based on the gold rules of prediction, the AltProt can present other domain and function actually undiscovered. If some AltProt can show a sequence homology with protein find for the moment specific to other species than Human, so they can explain reaction to human at other virus origin. Also probably pathway missing link can be complete by the AltProt studies.

The experiments carried out in this study make it possible to demonstrate the interactions of viral proteins with the proteins of the host cell. From this context we have no information on the protein interactions inside of the host cell, so the determination of the functions of the identified AltProts is difficult, since the identified AltProts can be linked to all the signaling pathways affected by the viral protein. Domain homology allows us to speculate on the function of these of the 27 AltProts with homology. For the others (32 proteins with <80% homology or without homology) considering their viral interacting protein and the RefProts that interact with these viral proteins, it is possible to hypothesize the signaling pathways involving these AltProts. In this way, among the five AltProts interacting with the viral protein “E”: IP_219869 (AltDGKH), IP_724315 (AltHMGN2P3), IP_788706 (AltEIF2S2P3), IP_555327 (AltAC006386.1) & IP_594707 (AltEEF1A1), three do not present an homology up to 80% with a RefProt (IP_219869, IP_555327, IP_594707), however the study of Gene Onthology of RefProts found in interaction with E (Fig.2-B), shows that the most represented Biological Processes are: "regulation of histone H3-K36 trimethylation" and “Synaptic vesicle budding from endosome” represented by the presence of RefProt: BRD4 and AP3B1. Thus these three AltProt, all like IP_724315 (AltHMGN2P3) homologous to the “non-histone chromosomal protein HMG-17”, may be involved in modifications of histones or the chromosomal binding and therefore in epigenetic phenomena.
In the same way six AltProts interact with the viral protein "M", among them two do not present any homology with RefProts. However the other four are homologous with kind of Tubulins, moreover the Gene Ontology analysis of RefProts in interaction with M (Fig.2-C), presents the main Biological Process: "microtubule nucleation by microtubule organizing center". It’s safe bet that the two AltProts of unknown function are involved in microtubule organization and protein transport. Finally the two AltProts (IP_671071, IP_565887) exhibiting low homology with bat proteins and observed in interaction with Orf8, can be proteins from the cytoskeleton such as the AltProts IP_774695, IP_593099, IP_774693 and IP_656465 exhibiting strong homologies with the tubulin family, but may also be linked to the post-translational glycosylation modification signaling pathway, such as the Biological Processes of RefProts interacting with Orf8 (Fig.2-D).

Further work is still needed to establish a complete and validated database of SARS-CoV-2 proteins interactome including AltProts and then further specify the exact role of these AltProts in pathophysiological mechanism of the viral infection. Nevertheless, some AltProts are already foreseen to be key player in the virus-cell hijacking, such as AltHSPA8P11, which is found to interact with 7 viral proteins. A cluster of AltProts centered on nsp6, nsp10, nsp11, Orf3b, Orf6, Orf7a and Orf9b is also identified. Very interestingly, most of these proteins are involved in the interferon production inhibition, innate immunity modulation, cycle arrest and host translation inhibition21. A major interest of the large scale interactomics is the possibility to screen for drug repurposing, as presented by the authors in their initial study. AltProts must now be considered as new potential therapeutic targets. Indeed, among the AltProts identified, the IP_2336782 (AltDUSP4) is found to be in interaction with Nsp6. AltDUSP4 shares 54% sequence homology with the C3a anaphylatoxin chemotactic receptor (C3AR1), which was recently shown to be involved in severe forms of COVID-19. C3AR1 is found over-activated in some patients leading to an hyper-inflammatory profile inducing persistence of the virus and a strong immunopathology [31]. Thus, AltDUSP4 is a potential target to reduce severe symptoms of COVID-19. Interestingly, the search for partner molecules via IUPHAR/BPS Guide to Pharmacology and BindingDB, shows the presence of sequence similarity between AltDUSP4 and the ATP binding cassette subfamily G member 2. It has to be noted that the viral protein Nsp6 was previously identified as a target of Bafilomycin A1, a potent and selective inhibitor of the vacuolar H+-ATPase. Several drugs are known to be active towards ATPase activity e.g. like cyclosporin A; KS 176; compound 14; Ko143 and Fumitremorgin C. and thus can target both NSP6 and AltDUSP4.

Taken together, these new findings highlight the presence of many yet unknown proteins in the interactome between the host cells and the viral proteins which are involved in major pathways such as innate immune response or translation regulation. This establish that besides the reference proteome, a ghost proteome exists, whose consideration would be highly beneficial both to the understanding of pathophysiological mechanism of the virus and to establish the therapeutic strategies.
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DECLARATION OF INTERESTS
The authors declare no competing interests.

AUTHORS CONTRIBUTION
Conceptualization: MS, IF; Methodology: TC; Formal Analysis: TC; Investigation: MS, TC; Resources: MS, IF; Data curation: TC; Writing: IF, MS, TC; Original Draft: IF, MS, TC; Supervision, Project: IF, MS, TC; Administration: MS, IF; Funding Acquisition: IF, MS

EXTENDED DATA
Table. List of AltProts identified to be interacting with the SARS-Cov-2 viral proteins
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Figure 1: Viral protein interaction network (SARS-Cov-2) from Co-IP experiments [21], re-analyzed by including AltProt, RefProt and viral databases. The previously established network is compared to the new query thanks to the DyNet Analyzer application on Cytoscape V3.8.0. Color legend nodes: red: viral protein (bait), blue: RefProts and green: AltProts, and for the edges: red: interaction not recovered in our analysis, grey are recovered in both analysis, green: specific to our analysis and with a ratio<100, purple edges are interaction specific to our analysis with a ratio of 100.
Figure 2: Gene Ontology analysis based on the RefProt identified in the network of interaction. ToppGene analysis is performed A- based on the gene name of the RefProt identified to have more than 80% of homology and on the RefProt identified in interaction with the bait of the CoIP, for B- RefProt in interaction with E, C- in interaction with M and D- in interaction with Orf8. The pathway attributed to the RefProt is a clue for the AltProt link to the same bait.
Table 1: List of AltProts identified to be interacting with the SARS-Cov-2 viral proteins. Co-IP raw data were re-interrogated using OpenProt [18]. The table lists the 56 identified AltProts identified including the name of the gene coding for the RNA transcript, the accession number of the transcript, the name of the AltProt, the type of transcript for which AltProts are issued from and for AltProts originating from mRNA, the location on the mRNA.

| Accession | GN               | TA                | AltProt   | type     | location |
|-----------|------------------|-------------------|-----------|----------|----------|
| IP_555327 | AC006386.1       | ENST00000624491   | AltAC006386.1 | mRNA     | 3'UTR    |
| IP_581922 | AC018641.7       | ENST00000435950   | AltAC018641.7 | ncRNA    | -        |
| IP_659614 | ANKRD20A11P      | ENST00000442192   | AltANKRD20A11P | ncRNA    | -        |
| IP_187691 | ARL3             | NM_004311.3       | AltARL3   | mRNA     | 3'UTR    |
| IP_077449 | BATF3            | NM_018664.2       | AltBATF3  | mRNA     | CDS      |
| IP_2387661| BCL11A           | XM_017004337.1    | AltBCL11A | mRNA     | 5'UTR    |
| IP_565887 | C9orf116         | ENST00000371789   | AltC9orf116 | mRNA     | 5'UTR    |
| IP_075271 | CDC73            | XM_006711537.3    | AltCDC73  | mRNA     | CDS      |
| IP_766056 | CEP290           | ENST00000547691   | AltCEP290 | mRNA     | 5'UTR    |
| IP_691726 | CTC-398G3.1      | ENST00000483614   | AltCTC-398G3.1 | ncRNA    | -        |
| IP_219869 | DGKH             | NM_152910.5       | AltDGKH   | mRNA     | CDS      |
| IP_2336782| DUSP4            | XM_011544428.2    | AltDUSP4  | mRNA     | 5'UTR    |
| IP_235699 | EDC3             | ENST00000565602   | AltEDC3   | mRNA     | 3'UTR    |
| IP_594707 | EEF1A1           | ENST00000309268   | AltEEF1A1 | mRNA     | 5'UTR    |
| IP_788706 | EIF2S2P3         | ENST00000428356   | AltEIF2S2P3 | ncRNA    | -        |
| IP_2396759| GJA5             | XM_017001044.1    | AltGJA5   | mRNA     | 5'UTR    |
| IP_711582 | HGS              | ENST00000577012   | AltHGS    | mRNA     | 5'UTR    |
| IP_775502 | HIGD1AP10        | ENST00000527837   | AltHIGD1AP10 | ncRNA    | -        |
| IP_724315 | HMGN2P3          | ENST00000433603   | AltHMGN2P3 | ncRNA    | -        |
| IP_557348 | HNRNPA1P28       | ENST00000424481   | AltHNRNPA1P28 | ncRNA    | -        |
| IP_572435 | HSPA8P11         | ENST00000508840   | AltHSPA8P11 | ncRNA    | -        |
| IP_658154 | HSPD1P7          | ENST00000447985   | AltHSPD1P7 | ncRNA    | -        |
| IP_289249 | KCNE1            | XM_017028342.1    | AltKCNE1  | mRNA     | 3'UTR    |
| IP_075761 | LAD1             | ENST00000631576   | AltLAD1   | mRNA     | CDS      |
| IP_671071 | LOC101929023     | ENST00000434879   | AltLOC101929023 | ncRNA    | -        |
| IP_2361135| LOC102723525     | XR_0925379.2      | AltLOC102723525 | ncRNA    | -        |
| IP_2268667| LOC105372714     | XM_017028195.1    | AltLOC105372714 | mRNA     | 5'UTR    |
| IP_2266298| LOC107985441     | XR_001754616.1    | AltLOC107985441 | ncRNA    | -        |
| IP_2354489| LOC107986350     | XR_001742414.1    | AltLOC107986350 | ncRNA    | -        |
| IP_143572 | LYRM2            | NM_020466.4       | AltLYRM2  | mRNA     | 3'UTR    |
| IP_745252 | MEG8             | ENST00000553465   | AltMEG8   | ncRNA    | -        |
| IP_213668 | METAP2           | XM_005268583.3    | AltMETAP2 | mRNA     | CDS      |
| IP_729791 | MT1X     | ENST00000568370 | AltMT1X | mRNA   | 3'UTR  |
| IP_230046 | NKK2-1-AS1 | ENST00000521292 | AltNKK2-1-AS1 | ncRNA | -     |
| IP_597201 | NOP56P1  | ENST00000440030 | AltNOP56P1 | ncRNA | -     |
| IP_105102 | POCTA    | XM_0115333561.1 | AltPOCTA | mRNA   | CDS   |
| IP_581419 | RP11-10F11.4 | ENST00000634439 | AltRP11-10F11.4 | ncRNA | -     |
| IP_734708 | RP11-24M17.3 | ENST00000567565 | AltRP11-24M17.3 | ncRNA | -     |
| IP_667059 | RP11-397P13.7 | ENST00000427282 | AltRP11-397P13.7 | ncRNA | -     |
| IP_591742 | RP11-471B18.1 | ENST00000407538 | AltRP11-471B18.1 | ncRNA | -     |
| IP_612631 | RP11-553P9.1 | ENST00000509116 | AltRP11-553P9.1 | ncRNA | -     |
| IP_639311 | RPL36AP13 | ENST00000457490 | AltRPL36AP13 | ncRNA | -     |
| IP_750273 | RPL4P1   | ENST00000496596 | AltRPL4P1 | ncRNA | -     |
| IP_637436 | RPL5P9   | ENST00000448118 | AltRPL5P9 | ncRNA | -     |
| IP_597129 | RPS17P1  | ENST00000396783 | AltRPS17P1 | ncRNA | -     |
| IP_668819 | RPS23P9  | ENST00000448848 | AltRPS23P9 | ncRNA | -     |
| IP_594653 | SENP6    | ENST00000474906 | AltSENP6 | ncRNA | -     |
| IP_769089 | SPRYD4   | ENST00000338146 | AltSENP6 | ncRNA | -     |
| IP_713094 | SSTR2    | ENST00000357585 | AltSSTR2 | mRNA   | 3'UTR |
| IP_656465 | TUBA3GP  | ENST00000410028 | AltTUBA3GP | ncRNA | -     |
| IP_774695 | TUBAP2   | ENST00000530835 | AltTUBAP2 | ncRNA | -     |
| IP_774693 | TUBAP2   | ENST00000530835 | AltTUBAP2 | ncRNA | -     |
| IP_593099 | TUBB2BP1 | ENST00000404155 | AltTUBB2BP1 | ncRNA | -     |
| IP_572422 | TUBBP1   | ENST00000518096 | AltTUBBP1 | ncRNA | -     |
| IP_665452 | UBE2D3P1 | ENST00000436669 | AltUBE2D3P1 | ncRNA | -     |
| IP_274314 | ZNF569   | XM_006723046.2  | AltZNF569 | mRNA   | 3'UTR |

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