RESEARCH ARTICLE

Alternative splice variants of rhomboid proteins: Comparative analysis of database entries for select model organisms and validation of functional potential [version 2; referees: 1 approved, 1 approved with reservations, 1 not approved]

Previously titled: Alternative splice variants of rhomboid proteins: In silico analysis of database entries for select model organisms and validation of functional potential

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Abstract

Background: Rhomboid serine proteases are present across many species and are often encoded in each species by more than one predicted gene. Based on protein sequence comparisons, rhomboids can be differentiated into groups - secretases, presenilin-like associated rhomboid-like (PARL) proteases, iRhoms, and “inactive” rhomboid proteins. Although these rhomboid groups are distinct, the different types can operate simultaneously. Studies in Arabidopsis showed that the number of rhomboid proteins working simultaneously can be further diversified by alternative splicing. This phenomenon was confirmed for the Arabidopsis plastid rhomboid proteins At1g25290 and At1g74130. Although alternative splicing was determined to be a significant mechanism for diversifying these two Arabidopsis plastid rhomboids, there has yet to be an assessment as to whether this mechanism extends to other rhomboids and to other species.

Methods: We thus conducted a comparative analysis of select databases to determine if the alternative splicing mechanism observed for the two Arabidopsis plastid rhomboids was utilized in other species to expand the repertoire of rhomboid proteins. To help verify the in silico observations, select splice variants from different groups were tested for activity using transgenic- and additive-based assays. These assays aimed to uncover evidence that the selected splice variants display capacities to influence processes like antimicrobial sensitivity.

Results: A comparison of database entries of six widely used eukaryotic experimental models (human, mouse, Arabidopsis, Drosophila, nematode, and yeast) revealed robust usage of alternative splicing to diversify rhomboid protein structure across the various motifs or regions, especially in human, mouse and Arabidopsis. Subsequent validation studies uncover evidence that the splice variants selected for testing displayed functionality in the different activity assays.
Conclusions: The combined results support the hypothesis that alternative splicing is likely used to diversify and expand rhomboid protein functionality, and this potentially occurred across the various motifs or regions of the protein.

Keywords
Rhomboid proteins, alternative splicing, activity, transgenic expression, recombinant rhomboid proteins, protein structure, model organisms, databases

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Competing interests: Parts of this study may be considered as being potentially related to material contained in a patent application (US 2016/0129078 A1). There are no other competing interests disclosed at the time of submission.

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Amendments from Version 1

The reviewers’ feedback for version 1 was valuable in guiding our revisions. Version 1 was revised substantially to address the aspects raised and to enhance clarity for the readers in the areas of writing style and additional details.

The presentation of the study’s purpose and its approaches was revised for clarity. This was achieved first by removing most of the references listed from 2 to 26 so that the references cited were more obviously directed at the study, as opposed to providing a general background of the wider field. Accompanying revisions were then made to the Abstract and Introduction.

The Methods section was revised substantially to clarify the approaches used and to provide more needed details concerning the comparative analysis of database entries and the functionality assays using recombinant proteins. The title was subsequently modified slightly to align with these revisions. The Results and Discussion sections were modified similarly to reflect the changes. The multi-year facet of the database analysis was deemed unnecessary and removed. Revisions were also made to the Source Datasets and the Supplementary Tables to enhance their presentation. Collectively, these revisions should help the readers interpret the work more clearly with the additional details and with the streamlining of the protocol for the database work.

See referee reports

Introduction

Rhomboid proteins are found widely in all types of organisms, spanning bacteria, archaea, eukaryotes. In higher-order organisms, rhomboid proteins are often encoded by a large group of genes, for example, upwards of twenty-two database entries reported for Arabidopsis and thirteen for humans (assessed as of May 2017). Phylogenetic studies, such as that conducted by Lemberg and Freeman (2007), suggest that rhomboid genes can be divided into two subgroups encoding proteolytically active secretases- and Presenilin Associated Rhomboid Like (PARL)-type rhomboid forms, and two subgroups of catalytically inactive forms (“non-proteases”) such as iRhoms, Derlins, and other distantly related forms.

The ‘active’ category of structurally diverse rhomboid proteases are often found to occupy regulatory roles of different cellular activities, typically by interacting with and cleaving specific membrane-residing substrates. Examples being the epidermal growth factor signaling pathway in fruit flies, the quorum-sensing mechanism of the bacterium Providencia stuartii, yeast mitochondrial membrane remodelling, and the health status of mitochondria in human cell lines.

The ‘inactive’ subcategories consist of structurally diverse rhomboid proteins as well and are believed to lack the needed catalytic residues used by the active rhomboid proteases. Despite the absence of the catalytic residues, some of the inactive rhomboid proteins are found to be functionally significant without being active proteases. In some of the reported cases, interaction alone with an inactive rhomboid, without proteolysis, is sufficient to cause effects, such as growth signaling in cancer cells (human iRhom1), dislocation of proteins in the endoplasmic reticulum protein (ER) (mammalian Derlins), ER protein quality control (Drosophila iRhom Rhomboid-5), development, and organelle biogenesis (Arabidopsis At1g74130).

In addition to the plethora of already functionally diverse active and inactive rhomboid proteases, alternative splicing appears to generate even more structural variants with modified functionalities. For example, one case was confirmed for the human RHBD2 gene. Levels of the two alternatively spliced RHBD2 mRNAs were elevated in breast cancer cell lines, suggesting a link between cancer cell activity, and the presence of splice variants with a sizable difference between the two variant protein structures. In Arabidopsis, the active plastid rhomboid At1g25290 was confirmed to exist as two functionally significant splice variants that differ by the presence of a potential cyclin-binding motif, a motif known to be involved in cell cycling.

To date, the phenomenon of diversifying rhomboid protein functionality through alternative splicing has been documented for two Arabidopsis plastid rhomboids. It has not yet been assessed as to whether this phenomenon is limited, or represents a mechanism for expanding the number of functional rhomboid forms in a wide range of rhomboid systems and organisms. Therefore, using the findings reported for At1g25290 and At1g74130 as guidance, we conducted a comparative analysis of entries available in the genetic databases of six widely used eukaryotic experimental models to assess how splice variants may be reflective of ways to diversify functionality, from limited amino acid changes to substantive structural deletions. A limited selection of alternatively spliced variants were then analyzed to document evidence of potential activity as proteins using different types of assays.

Methods

Comparison of variant sequences in current databases and categorization scheme

The sequence entries compared were assembled from current versions of the publicly-accessible databases (last sampled as recent as May 31, 2017). We used the NCBI database (RefSeq, RRID:SCR_003496) as our primary source for the six widely-used model organisms (Homo sapiens (human), Mus musculus (mouse), Arabidopsis thaliana (Arabidopsis), Drosophila melanogaster (fruit fly), Caenorhabditis elegans (nematode) and
Saccharomyces cerevisiae (Baker’s yeast). We also cross-checked with other databases such as the Mouse Genome Informatics resource (Mouse Genome Informatics, RRID:SCR_006460), TAIR for Arabidopsis (TAIR, RRID:SCR_004618), FlyBase (FlyBase, RRID:SCR_006549), WormBase (WormBase, RRID: SCR_003098), and the Saccharomyces Genome Database (SGD, RRID:SCR_004694). All entries used and assessed are compiled in Table S1 along with their relevant details. Note that some of the entries listed in supplementary table are for the predicted forms, despite being assessed by comparison to RNA-seq data, for example. Table S2 provides a listing of formal and alternate names used for each rhomboid, along with Gene ID numbers and related references. Database entries were compiled and assessed for alternative splicing before use by comparing entries using publicly available bioinformatic tools for alignment work (the suite of BLAST (RRID:SCR_007190), Clustal (RRID: SCR_001591), and LALIGN tools). Entries were retrieved from databases for cDNA/EST sequences, RNA-seq, genomic sequences, and proteins. This assessment stage was not designed to correct predictions, address gaps, or add predictions, but to assess the existing entries and the overall capacity of alternative splicing to diversify rhomboid protein functionality. This study’s objective was thus focused on surveying the functional potential of the available entries with the understanding that some will require further validation work. The categorization of alternative splicing events/splice variants was based on the potential impact of changes on motifs along the protein, from amino to carboxyl terminus. The use of six eukaryotic rhomboid gene systems was to help determine common trends. The protein models and motifs used for categorization were adapted from the models reported by Lemberg and Freeman. In many cases, alternative splicing events tend to influence one motif, but there were a number of cases where the impact may affect one or more motifs, depending on the type and location of the splicing event. For the purpose of comparing the impacts of alternative splicing, all variant sequences were assumed to result in the generation of functional proteins in some capacity, including extensively truncated products. For this study, sequences and splicing events are categorized at the motif-specific or region-specific level and may thus appear in more than one category. For instance, a frequent occurrence is an alternative splicing event which occurs in a particular motif that would likely impact the adjacent linker region. The various splice variants are listed in Tables S3 to S10 by category. Information concerning the impact of the splicing events is also included in these Supplementary Tables. Overviews are provided in the Results section. Specific examples are highlighted when applicable.

Alignments and analyses of changes to protein structure
Linear protein alignments were carried out first using the publicly available bioinformatic tools listed in the above section, such as Clustal Omega (RRID:SCR_001591). Select sets of the linear alignments are included in the Supplementary Material. Structural predictions were facilitated by comparing alignments with predicted structures constructed and reported by Lemberg and Freeman. Three-dimensional predictions of splice variant protein structures were created and compared by utilizing Phyre Version 2 services (RRID:SCR_010270) and visualized using PyMol 1.1 (RRID:SCR_00305). Comparisons were conducted using the bacterial rhomboid GlpG model. These 3D predictions were carried out solely for the purpose of investigating and speculating the capacity for structural impact of the different splicing events. All of these 3D predictions are provided and reported in the Supplementary Material for consideration only and do not represent established, solved structure data.

Production of select splice rhomboid protein variants and immunoblot assessments
Select proteins were synthesized in Escherichia coli JM109 (DE3) and facilitated by the T7 promoter of pET20b (EMD4 Biosciences, Rockland, MA, USA). Histidine tags were joined in-frame to the carboxyl-terminus. Cells were grown at 16°C in ampicillin-containing Terrific Broth (Bioshops Inc., Burlington, Canada) (25 µg/ml ampicillin). Recombinant histidine-tagged proteins were purified using nickel – nitriolaecetic acid affinity chromatography (Qiagen, Toronto, Canada). The protein expression and chromatography procedures followed and the composition of the buffers used were as reported and cited previously. Briefly, bacterial cells were harvested by centrifugation, resuspended in a small volume of extraction buffer, and disrupted using a French Pressure Cell Press (2 cycles of 15,000 psi using a medium cell). Elution was carried out in one step using a small volume of the cited pH 7.5 elution buffer and 400 mM imidazole. Proteins were quantified using the Bradford assay system (Biorad, Hercules, CA, USA) and normalized before use. Proteins were typically stored as concentrated stocks at -80°C and used later for the various assays. Freeze-drying of recombinant rhomboid proteins did not have an observable impact on activity.

Protein samples, typically 0.5 µg per lane, were analyzed when needed using standard one-dimensional (1D) 12% (m/v) sodium dodecyl sulfate (SDS) – polyacrylamide gels. Electrophoretic and immunoblotting protocols were performed according to Laemmli and Towbin. Immunoreactive bands were analyzed using scans, quantitated by densitometry (Scion Image 4.0.3.2, Scion Corporation, USA), normalized, and compared relative to internal references when applicable. All immunoblots were repeated at least three times within each experiment and with biological replicates. Representative results are then shown in the figures. When needed, quantitations were conducted using nonsaturated scans of the images presented in the figures. Various recombinant protein preparations were checked by immunoblotting using rabbit polyclonal anti-rhomboid protein antibodies that were established by our lab and validated previously as reported in Powles et al. For samples derived from transgenic yeast cell assays, rabbit polyclonal anti-yeast mHsp70 antibodies (Antibodies-Online Cat# ABIN488515, RRID:AB_11209968) were used for normalization as reported previously in Powles et al. The normalized ratio strategy was designed to allow semi-quantitative assessment of profile changes independent of experiment, gel origin, and exposures. Statistical analyses or details are noted where applicable.

Assays for testing biological activity of select splice rhomboid protein variants
Activity assays using transgenic yeast – Protein expression in S. cerevisiae (Baker’s yeast) was facilitated by the yeast – E. coli shuttle vector pACT2 (BD Biosciences-Clontech, San Jose, CA,
USA). In all cases, expression of the inserted cDNAs was driven by the cloned yeast Rbd1 promoter. The introduction of plasmids was carried out using standard yeast transformation techniques. The host yeast strain C6000 used was acquired from EURO- SCARF (EUROpean Saccharomyces Cerevisiae ARChive for Functional Analysis, RRID:SCR_003093) (Frankfurt, Germany). Cells were grown in glucose-supplemented media (at 30°C in standard glucose-supplemented yeast Complete Medium without leucine where applicable (20 g/l glucose, 6.8 g/l yeast nitrogen base without amino acids, (Sigma-Aldrich, Oakville, ON, Canada), 1.6 g/l drop out mix (Sigma-Aldrich, Oakville, ON, Canada), and 20 g/l agar when used). All strains were prepared as populations (as opposed to from single colonies) and stored at -80°C as glycerol stocks.

A disk diffusion method was employed to test different yeast strains for changes to nystatin sensitivity as a result of the indicated splice variant proteins being expressed. Yeast cells were suspended in top agar (1% w/v) media at 5 × 10⁶ cells/mL and poured onto agar plates of the same medium used above (2% w/v). Sterilized filter paper disks (38.5 mm²) infused with nystatin, were evenly positioned on the top agar, at a typical concentration of 10 disks per 90 mm plate. Each disk was infused with 10 µL of a 2.4% (v/v) nystatin solution, diluted with the appropriate yeast culturing medium. Plates were incubated overnight at 30°C. The zone of growth inhibition around each disk was then measured, and the corresponding area calculated. Multiple independent experiments were conducted to confirm the functional potential of the various splice variants.

Activity assays using yeast and externally-added proteins - Non-transformed cells (without plasmids) were also assessed with exogenously added splice variant proteins. Since amphotericin B (AmB) has the ability to introduce transmembrane channels for protein delivery, cells were incubated for one hour with 10 µg (at 1 µg/mL, made in 23X diluted elution buffer (diluted with growth media) of the indicated recombinant variant protein along with a 1% (v/v) AmB solution (this level has no detrimental effect on yeast cell survival). Controls were carried out prior to the experimental assays with various diluted elution buffer levels to assess the base level effects without recombinant rhomboid proteins. When shown, the controls or mock assays were conducted with the above 23X diluted elution buffer. Minimal volumes of cells, typically at 5,000 cells/mL, were used. Various levels of nystatin were then added and incubated for the last 15 minutes of the 1 hour treatment period. After the 1 hour incubation-treatment, 500 cells were plated (per 90 mm petri plate) and grown for 48 hours at 30°C to assess sensitivity to nystatin as a test for biological activity and for any differences between splice variants. Results from independent replicates were analyzed statistically (t-test) between control (or mock) and experimental assays and noted with details where applicable.

Activity assays using transgenic bacteria – E. coli JM109 (DE3) cells harboring various pET20b-based splice variant constructs were plated on LB agar containing varying concentrations of ampicillin as indicated. The assessment of ampicillin sensitivity was not dependent on induction of expression, but instead relied upon the inherent “leaky” expression. Plating was normalized with equal colony numbers. Results from independent experiments were analyzed statistically (t-test) between control (or mock) and experimental assays, and noted with details where applicable. Changes in ampicillin sensitivity were further examined using whole cell extracts and immunoblotting to assess β-lactamase expression, secretion, and processing of the precursor β-lactamase form. Whole cell extracts were prepared from cell pellets harvested by centrifugation of liquid cultures and boiled in standard protein gel loading dye.

Activity assays using bacteria and externally added proteins - Bacteria (HB101) were assessed in some cases with exogenously added splice variant proteins. Cells harboring pET20b were used as the ampicillin resistance model for testing biological activity and differences between splice variant proteins. Dimethyl sulfoxide (DMSO) was utilized to permit the delivery of proteins into cells by an initial incubation of 30 minutes with 5% (v/v) DMSO, 10 µg of variant protein (normalized with 23X diluted elution buffer (diluted with LB broth)), and LB broth. The controls used here for bacterial cells were established and conducted in the same manner as described for the above yeast cell assays. After the 30 minute protein delivery period, each treatment was incubated with 1.25 or 1.5 mg/mL of ampicillin (and adjusted if needed) for an additional 45 minutes. Treatments were carried out at 37°C with shaking (100 rpm). Bacteria were normalized to 800 cells per treatment in a total volume of 200 µL, and plated in its entirety on LB plates and grown overnight at 37°C. Surviving colonies were counted and compared to mock treatments, which consisted of all components without proteins. Results from independent experiments were analyzed statistically (t-test) between control (or mock) and experimental assays and noted with details where applicable.

Results

Rationale and justification for this database study

We previously verified in separate studies a mechanism for diversifying rhomboid proteins and their functionality. This alternative splicing mechanism played diversifying roles for two different Arabidopsis plastid rhomboid genes - the active secre- tase type At1g25290 and the inactive PARL type At1g74130. Alternative splicing impacted different parts of the proteins with no apparent functional similarities to each other. The At1g25290 splice variants were focused on controlling the appearance of the cyclin-binding RVL motif in the protein’s middle segment, right after the third predicted transmembrane region. The data underlying the characterization of the At1g25290 splice variants (designated L and S) and the splicing events involved to generate the protein variants were reported previously. The composition of the resulting At1g25290 protein variants (L and S) used later in this study are shown in the Supplementary Material. The splice variants created for At1g74130 resulted in different shortened proteins, each missing a key glutamine residue in the last carboxyl transmembrane region. Alternative splicing resulted in a substantial impact to the carboxyl transmembrane segment of At1g74130, changing from a seven predicted to six transmembrane structure. The data underlying the characterization of the At1g74130 splice variants (designated L, M, and S) and

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the splicing events involved to generate the protein variants were reported previously\textsuperscript{23,24}. The composition of the resulting At1g74130 protein variants (L, M, and S) used later in this study are also shown in the Supplementary Material. Both studies provided evidence that the resulting variant proteins display altered functionality\textsuperscript{23,24}. These two sets of findings alone bring the total number of plastid rhomboid forms in \textit{Arabidopsis} to at least seven, two for At1g25290, three for At1g74130 and at least one each for At1g74140 and At5g25752.

The outcomes discussed above prompted us to look at rhomboid genes of other eukaryotic species for evidence of similar diversification mechanisms. We thus compared and analyzed the RNA sequence databases of six eukaryotic organisms used widely as experimental models - human, mouse, \textit{Arabidopsis}, \textit{Drosophila}, \textit{C. elegans}, and \textit{S. cerevisiae}. Even though the databases continue to evolve, the observations disclosed here should continue to be applicable.

Is alternative splicing present in different rhomboid gene systems?
The first aspect to establish was the presence of alternative splicing and its extent within a selected eukaryotic species, and across the different selected species. Using the current versions of the RNA sequence databases, we compiled and assessed all possible RNA sequence entries that were derived from alternative splicing. The human, mouse, and \textit{Arabidopsis} assessments revealed many potential alternative splicing events for different rhomboid genes of these species. There were 95 entries in human, 53 in mouse and 40 in \textit{Arabidopsis} (Figure 1). In contrast, similar analyses of \textit{Drosophila}, \textit{C. elegans} and \textit{S. cerevisiae}, revealed minimal levels to no evidence of alternative splicing. We found one possibility each for \textit{Drosophila} and \textit{C. elegans} and none so far for \textit{S. cerevisiae}. It is, however, possible that the outcomes observed for the latter three model species were due to the number of reported alternate RNA sequences at the time of assessment. This was the case in \textit{Arabidopsis} where alternative splice variants for At1g25290 and At1g74130 were discovered and verified upon further analysis of transcript populations\textsuperscript{23,24}.

Of the six eukaryotic species analyzed, the human rhomboid system appears to exhibit the most alternative splice variants. All 13 of the rhomboid or like genes display multiple entries reflective of alternative splicing (see Figure 2 and Table S1). For instance, human PARL contains verified splice variants and additional predicted mRNA sequences or proteins. Similar situations were observed for human RHBDL2 (iRhom2), RHBDL1, RHBDD1, and RHBDD2.

The use of alternative splicing is also evident in the mouse rhomboid genes. The assessment revealed evidence of multiple alternative splice variants for mouse rhomboids (Figure 1 and Table S1).

The relatively small genome of \textit{Arabidopsis} appears to possess a high number of rhomboid and like genes. There are 22 entries and 10 are accompanied by 1 or 2 additional splice variant sequences (Figure 1 and Table S1). The splice variants arising from At1g25290 and At1g74130 were discovered and verified in two other studies\textsuperscript{23,24}. Based on the trends observed for human and mouse, it is likely that there are other splice variants in \textit{Arabidopsis} awaiting discovery, especially for the other 12 gene entries currently without accompanying sequence variants in the database.

What are the potential types of changes introduced by alternative splicing?
Further analyses of the Figure 1 entries indicate that many of the occurrences are likely reflective of mechanisms for diversifying functionality (Figure 2 and Table S3–Table S10). Structural changes were observed for both active rhomboid proteases (secretases and PARLs) and rhomboid-like proteins (inactive rhomboids and iRhoms) (Figure 3–Figure 6, Table S4–Table S9). Potentially impactful changes were located in domains

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**Figure 1.** Visual summary of available sequence entries retrieved and studied for the selected eukaryotic organisms. Each of the selected eukaryotic organism is indicated along with the total number of rhomboid and rhomboid-like genes found in the databases and the number of accompanying variant sequences obtained and studied in this report. Entry details are summarized in Table S1. All entry types were analyzed.
Figure 2. Schematic summary of splice variant types associated with different regions of a generalized rhomboid protein. Each region of the generalized rhomboid protein is indicated along with the total number of rhomboid and rhomboid-like genes that were accompanied by splice variants found in the databases. The genes considered belong only to the eukaryotic organisms selected for study here and are tallied together independent of species. The total number of accompanying alternative splice variants determined for each region is also provided. These variants are again tallied independent of species. Entry details are summarized in Table S1.

Figure 3. Locations of splice variants in the secretase-type rhomboids. Alternative splicing and resulting changes to the secretase-type rhomboid proteases (6+1 model). Amino acid sequences of the splice variant proteins were aligned and compared. The outcomes of these comparisons are compiled in the Supplementary Tables. The results are summarized in this figure. The regions of the protein potentially influenced by the changes are annotated. The protein structure is constructed using information from Lemberg and Freeman. Information concerning the names used in the figure is provided in Table S2.

Changes were also observed within the 5' UTRs that may affect translation and 3' UTRs that may affect transcript properties (Table S3 and Table S10). The changes impacting the protein can be subtle, affecting a few amino acid residues, to entire sections of the protein. In some instances, there were extensive deletions, insertions, truncations, or shortenings of the protein.
Figure 4. Locations of splice variants in the PARL-type rhomboids. Alternative splicing and resulting changes to the PARL-type rhomboid proteases. Amino acid sequences of the splice variant proteins were aligned and compared. The outcomes of these comparisons are compiled in the Supplementary Tables. The results are summarized in this figure. The regions of the protein potentially influenced by the changes are annotated. The protein structure is constructed using information from Lemberg and Freeman1. Information concerning the names used in the figure is provided in Table S2.

Figure 5. Locations of splice variants in the category of iRhom proteins. Alternative splicing and resulting changes to iRhom proteins. Amino acid sequences of the splice variant proteins were aligned and compared. The outcomes of these comparisons are compiled in the Supplementary Tables. The results are in this figure. The regions of the protein potentially influenced by the changes are annotated. The protein structure is constructed using information from Lemberg and Freeman1. Information concerning the names used in the figure is provided in Table S2.

Changes to the 5’ untranslated region of the transcript - One of the most widely reported alternative splicing mechanisms is designed to control entry into translation or protein translation itself11. This continues to be the case for the rhomboid genes examined here (Table S3). Alternative splice variants with potential effects on translation were found in human, mouse and Arabidopsis. Twelve of the 13 human, and nine of the 13 mouse rhomboid genes were accompanied by entries with changes to the 5’ UTRs. Interestingly, despite the higher number of genes documented for Arabidopsis, 5’ UTR splice variants were found only for 7 of the 22 rhomboid genes. However, the absence of evidence for the other 15 Arabidopsis rhomboid genes may be due to unreported sequences or awaiting discovery, which was the case with At1g74130 and At1g2529023,24.

Changes to the amino terminal region - Alternative splicing events affecting the amino termini appear to be common as well in our set of RNA sequences (Figure 2–Figure 6 and Table S4). Changes affecting the region between the start methionine to the first predicted TMD are placed into this category which includes frame-shifts, alternate starting methionines, insertions and deletions. Changes in this region could affect functional aspects like protein targeting and transport, membrane insertion, assembly and topology, and assembly with complexes. With the exception of human
Figure 6. Locations of splice variants in the "inactive" category of rhomboid proteins. Alternative splicing and resulting changes to "inactive" rhomboid proteins (using the 1+6 model). Amino acid sequences of the splice variant proteins were aligned and compared. The outcomes of these comparisons are compiled in the Supplementary Tables. The results are in this figure. The regions of the protein potentially influenced by the changes are annotated. The protein structure is adapted from Lemberg and Freeman. Information concerning the names used in the figure is provided in Table S2.

RHBD1, all of the other 12 human rhomboid genes are accompanied by alternative splice variants impacting their amino end sequences. The situation is slightly different in the mouse rhomboid system where 9 of the 13 genes contain variants impacting the amino end. Based on the mouse database, Rhbdf2, Rhbdd1 and Rhbdd3 do not so far have any variants impacting this region. Interestingly, only 3 Arabidopsis rhomboid genes have entries with predicted alternative splicing within the amino region. Of the 5 Drosophila genes, Stet, has an altered amino terminus resulting in an alternative start methionine. Of the 5 C. elegans genes, one has an altered amino terminus due to alternative splicing. This ROM-4, displays a frame-shift that shortens the length of the region that could still possess functionality as with the case of At1g74130.

Changes to the L1 Loop region – The L1 region contains a conserved “loop” structure. This structure lies either side of transmembrane helices and plays a role in rhomboid protease activity. The L1 loop is also partially inserted into the membrane. Site-directed mutagenesis of the conserved L1 loop residues revealed evidence that this loop controls the way in which rhomboids interact with lipid membranes. Our analyses of the Figure 1 entries indicate that the L1 Loop region is likely subject to alternative splicing (Figure 2–Figure 6 and Table S5).

In human, splice variants involving the L1 loop accompany 9 different rhomboid genes. Generally, the outcomes of the variants range from deletion of residues from part of the L1 loop region, to altering a few residues at one end or the other of the structure.

A similar trend was found for 4 of the mouse rhomboid genes, Rhbdf1 (iRhom1), Rhbddd2, Derlin2, and Ubac2. Again, the two splice events resulted in the loss of residues from the L1 loop.

The Arabidopsis database contains splice variants of the L1 loop for two genes, RBL4 and RBL14. Like in human and mouse, alternative splicing resulted in the removal of residues or large sections of the L1 loop.

The C. elegans ROM-4 gene exhibits an alternate start methionine and a frame-shift within the L1 loop, giving rise to only the beginning part of the loop.

Changes to regions affecting other structural aspects – This category is defined as changes to the linker or other transmembrane regions (TMD) with no currently assigned functions, as opposed to the regions with distinct functions discussed above and below. Since changes to these regions, subtle or extensive, could potentially affect the other functional aspects of the protein itself, or its interactions, it would be important to assess these splicing events.

In human, 11 of the 13 rhomboid gene entries examined are accompanied by splice variants in regions of the protein that fall under this category. Examples being PARL variants lacking TMD3 and part of TMD4, or the amino end of TMD1 (see Figure 3–Figure 6 and Table S9).

Changes to neighbouring regions of the catalytic dyad - The catalytic sites of rhomboid proteases consist of residues
 contributed from two different transmembrane domains, TMD4 and TMD6, when using the “6+1” model. PARL and secretase-type catalytic residues are characterized by the amino acids GxSx – H. The catalytic residues of iRhoms are characterized by the residues GPxx – H.

There are currently 4 entries for human PARL rhomboids accompanied by splice variants that impact catalytic potential by altering the GASG or H sites through limited deletions, or extensions and the subsequent loss of the serine and glycine residues (GA SG to GA) (Figure 2–Figure 6, Table S6).

Both human iRhoms, RHBDF1 and RHBDF2 (iRhom1 and iRhom2), contain frame shifts and early terminations within the L1 loop. RHBDL1 has a predicted variant of potential for this rhomboid by using an alternate start methionine is accompanied by a splice variant that may alter the catalytic gene to highlight in this category is RBL 14 (At3g17611). RBL14 is maintained in this RBL3 variant. The last histidine is removed entirely. Although the Gate and the catalytic site changes associated with 3' UTR of rhomboid transcripts result in a frame shift within TMD3. Of the 13 human rhomboid genes with entries in our data set, 6 are accompanied by alterations to the Gating TMD. PARL, RHBDF1, RHBDL1, RHBDL3, DERL2 and DERL3 are accompanied by forms with altered TMD6 (based on the 1+6 model). The most common resulting event appears to be early termination of the protein.

The same splicing outcomes appear to be present in our set of entries for mouse, Arabidopsis and C. elegans rhomboids. Mouse Rhbdl3, Rhbdf1 and Rhbdf2 (iRhom1 and iRhom2) are accompanied by variants with deletions or early terminations caused by frame shifts. The Arabidopsis data set contains two genes with predicted alternative spliced sequences affecting the gating TMD region. Arabidopsis RBL1 possesses an insertion between the catalytic TMD4 and the linker to the Gate. The RBL3 variant is missing both the gating TMD5 and the catalytic TMD6. The C. elegans ROM-4 variant is also missing the gating TMD as a result of a frame shift.

Changes to the catalytic regions - Most of the carboxyl termini changes are due to frame shifts, giving rise to different carboxyl sequences. In human, PARL, RHBDF2, RHBDL1, RHBDD1, and Derlin3, all contain variants with different carboxyl ends. The situation is similar in mice where Rhbdf1, Rhbdf2 and Rhbdl3 are accompanied by variants with different carboxyl ends (Figure 2–Figure 6, Table S8).

In Arabidopsis, At1g74130 and At5g07250 variants have shortened carboxyl termini. The three alternative splice variants of At1g74130 lack the entire predicted carboxyl terminus. The introduction of an stop codon in TMD6 (or further upstream) resulted in the early termination of translation. The At5g07250 (RBL3) variant lacks TMD5 and TMD6, but the carboxyl terminus is restored with the removal of the first 4 residues of the predicted motif.

Changes to the 3’ UTR region of the transcript - There are also changes associated with 3’ UTR of rhomboid transcripts (Table S10). In our set of entries, there are 9 human genes with splice variants in the 3’ UTR. Some of the variants possess longer 3’ UTR sequences, whereas others exhibit shorter 3’ UTRs.

A similar situation is observed for mice where 5 genes are associated with variants containing altered 3’ UTRs. Rhbdl2, Rhbdf1 and Rhbdf2 variants contain extended 3’ UTRs, whereas Rhbdl3 variants exhibit shorter 3’ UTRs.

The Arabidopsis genes At1g74130, At3g17611 and At3g58460 are accompanied by one variant each with shortened 3’ UTRs. One other gene, At2g29050 (NM_001084504.1), is represented without a predicted 3’ UTR in the database.

C. elegans ROM-4 has an alternate starting methionine and a frame shift that results in the removal of the catalytic residues.
What are the potential impacts on the rhomboid protein structure upon alternative splicing?

The data compiled above suggest potentially impactful changes to the functionality of the affected proteins, but this speculation is limited to linear protein sequences and motifs. We were next interested in testing out possible functionality changes using currently available predictive tools for 3-D protein structures, despite this highly speculative assessment tool. To this end, we decided to use the established 3-D structure/model of the bacterial rhomboid GlpG to test the potential impact exerted by the various splice variant types. The GlpG model is the most established of the rhomboids and offers a more complete structure for this analysis. It should be noted that there are caveats associated with the use of the bacterial GlpG to assess other rhomboid types, but this analysis is strictly focused on how changes could theoretically impact such a rhomboid structure. Because these assessments are judged as being too speculative, the outcomes are provided only as Supplementary Material (Supplementary File 1 and Figure S1–Figure S8). These outcomes may be useful starting points for guiding future structural studies that validate the outcomes. The actual impacts to functionality and structure of particular alternatively spliced protein variants also need to be studied individually through experimentation. This notion was assessed here for a selection of splice variants using the different types of activity tests. These tests were devised mainly to uncover evidence of functionality in splice variant proteins, as opposed to assigning possible biological roles. Different contexts were used to assess functionality of the selected splice variants, contexts ranging from transgenic expression to assays using recombinant proteins as exogenous additives.

The first series of tests were conducted for the Arabidopsis At1g74130 splice variants to obtain verification of functionality in a heterologous transgenic setting. For At1g74130, a functional relationship between its splice variants and a known yeast mitochondrial rhomboid substrate, Mgm1, was initially discovered using transgenic yeast. As shown previously in Powles et al., each At1g74130 splice variant impacted the Mgm1 ratio (the amount of uncleaved (97 kDa) to cleaved (84 kDa)). The At1g74130 M and S splice variants individually reduced the ratio by about a third (from ratios of 0.67–0.7 to 0.45 for M and 0.39 for S). Since the At1g74130 splice variants exist simultaneously in their natural Arabidopsis context, we further assessed different combinations of the same splice variants to see if such combined interactions influence the Mgm1 ratio, an indicator of splice variant activity. Further adjustments to the ratios by the various combinations of splice variants would uncover additional evidence of interaction and functionality. The results in Figure 7D (Dataset 1) indicate that various variant combinations possess the ability to influence the Mgm1 ratio, mainly more uncleaved Mgm1 (triggering higher ratios, from 0.81–1.09, instead of the ratios observed for cells expressing one variant at a time). The pACT2 control mitochondria displayed a similar Mgm1 ratio to that observed in Powles et al., at 0.66 ±0.002 versus 0.70 ±0.002, respectively (Figure 7D and Dataset 1). Yeast cells expressing the pair At1g74130 (L) and (S) displayed a ratio of 1.05 ±0.02, cells expressing At1g74130 (L) and (M) exhibited a ratio of 0.81 ±0.004, and cells expressing At1g74130 (S) and (M) resulted in a ratio of 1.09 ±0.03 (Figure 7D and Dataset 1). The top set of ratios is provided and adapted from Powles et al., to contrast the differences between single variant and double variant expression. These results demonstrate that the splice variants exhibit functionality in this setting.

We next assessed activity by looking at changes in sensitivity to the fungicide nystatin. Changes to sensitivity was assessed in two ways, growth/survival of yeast cells around nystatin-infused disks as a longer treatment strategy and nystatin treatments of cell cultures as a transient strategy. As shown in Figure 7A and B (Dataset 1), the expression of At1g74130 variants in yeast (especially (S) and (M)) increased nystatin sensitivity and cell death. The pACT2 control yeast strain and the strain expressing At1g74130 (L) displayed similar average areas of clearance (between 40–50 mm²), whereas the strains expressing At1g74130 (S) and (M) frequently displayed larger clearance areas of approximately 60–70 mm² on average.

The disk assay results were reflected in the liquid culture assays. Yeasts expressing the same At1g74130 variants were treated transiently with nystatin and amphotericin B prior to plating. Compared to untreated cells, all strains experienced decreased survivability when treated with nystatin and/or amphotericin B (Figure 7C and Dataset 1). All cells displayed decreased survival when treated with amphotericin B. The pACT2 control was around 60.13 ±2.83% and the splice variant lines were around 74–76% (At1g74130 (L) line was at 75.91 ±0.83%, (S) line was at 75.49 ±5.99% and (M) line was at 74.26 ±2.04%). When treated with nystatin, all At1g74130 lines displayed decreased survivability relative to the control strain (pACT2 control line was at 30.82 ±4.85% compared to At1g74130 (L) line at 15.98 ±1.48%, (S) line at 15.07 ±0.77%, and (M) line at 19.52 ±6.25%). When treated with both amphotericin B and nystatin, the control line displayed 15.19 ±0.45% survival, whereas the splice variant lines were in the 6–12% range (At1g74130 (L) line at 6.62 ±1.20%, (S) line at 11.61 ±3.33%, and (M) line at 8.39 ±1.90%). Overall, with the exception of the amphotericin B alone treatment, cells expressing any of the three At1g74130 splice variants displayed lower survivability when treated transiently with fungicide. Both of the fungicide settings (panels A and B; Dataset 1), and then panel C; Dataset 1 indicate that the At1g74130 splice variants possess functionality. The phenomenon observed in transgenic yeast were also reflected in the transgenic bacteria setting, where β-lactamase expression and secretion are high and considered to represent a “Superbug” (antibiotic resistance) model. Enhanced sensitivity to ampicillin was observed in bacteria expressing the At1g74130 (L) and (S) splice variants relative to (M) (Figures 8A and B; Dataset 2, respectively). Bacteria expressing At1g74130 (L) and (S) exhibited higher levels of sensitivity at lower ampicillin concentrations relative to (M) in this context. At the protein level, bacteria expressing At1g74130 splice variants displayed reduced synthesis, processing and secretion of β-lactamase (Figure 8C). In cells with pET20b only, most of the β-lactamase were present in the mature form (29 kDa) and at high total cell levels. In contrast, bacteria expressing At1g74130 splice variants exhibited shifts toward the precursor form (31.5 kDa), with (L) being the most impacted (Figure 8C). Additionally,
there were lower levels of β-lactamase overall in these same cells that may further contribute to the higher levels of sensitivity (Figure 8C). These results indicate that the splice variants display functionality in this setting.

Evidence of functionality was also observed using an exogenous additive approach, where recombinant splice variant proteins were used to pre-treat cells before testing for changes to antimicrobial sensitivity (see Methods). This treatment scheme would be considered a transient strategy. For yeast cells, recombinant splice variants were delivered using a sub-lethal level of amphotericin B as the pre-treatment step before testing for sensitivity (survival) to nystatin. Even though amphotericin B is a fungicide (especially at higher levels), it was feasible to utilize amphotericin B at sub-lethal levels (1% (v/v)) for delivery purposes because this compound is capable of altering the permeability of fungal membranes.
Figure 8. Activity assays for At1g74130 splice variant proteins using transgenic bacteria and as exogenous additives to “Superbug” bacteria. (A) Bacterial cells expressing one of the three different splice variant proteins were grown on LB-ampicillin plates from 1.0 mg/mL to 1.75 mg/mL (labelled L, S, and M, as in the previous figures). (B) Graphical representation of the survival levels observed for each of the plates shown in panel A (reflected as the numbers of colonies growing). The labelling is the same as in panel A. (C) Expression of At1g74130 variant proteins in bacteria alters production and secretion of β-lactamase. Immunoblot images of β-lactamase protein samples are shown for different bacterial strains grown in different ampicillin concentrations. The lanes contain lysates from control cells containing pET20b only (labelled Control), and one of the At1g74130 variants, L, S, and M (labelled as Variant L, S, or M). The smaller-sized bands marked “M” represents the mature β-lactamase form (29 kDa) and the larger-sized bands marked “P” represents the precursor β-lactamase form (31.5 kDa). The full immunoblot image used is provided in Figure S10. (D) Recombinant At1g74130 variant proteins were tested for activity (enhanced sensitivity to ampicillin in this case) as exogenous additives in the same manner as that shown for yeast in Figure 7. Cells (resulting colonies) surviving the different treatments are depicted as Percent Survival in the graph. The error bars represent standard deviations (n=3). The Percent Survival was calculated relative to the No Treatment control cell numbers. “No treatment” represents the bacterial culture used (diluted to the prescribed cell number tested as described in Methods). The variant protein being tested is labelled as in the panel A. DMSO indicates the use of DMSO (5% v/v) as the delivery agent. The Mock Treatment contains all components used except with no recombinant proteins added. All treatments involve exposure to 1.25 mg/mL ampicillin.

and allow rhomboid proteins cellular access. The treatment matrix used for these yeast assays is shown in Figure 9A. The yeast strain tested here was the same parental host line used in the earlier assays. Sensitivity to nystatin was then tested at a level of 0.5% (v/v). Overall, treatment resulted in smaller colonies at the time of plate growth documentation (compare treatment 1 to treatments 2 to 9 in Figure 9B and C; Dataset 3). All control or mock-type treatments display higher survival percentages compared to the three recombinant rhomboid pre-treatments (Figures 9B and C; Dataset 3). Relative to the Positive Control (considered 100% survival), the amphotericin B only treatment (AmB Control) displayed 96.46 ±0.43% survival, nystatin only (Nys Control) showed 85.46 ±2.60%, the amphotericin B-nystatin treatment (AmB/Nys Control) was at 81.81 ±1.86%, the amphotericin B-nystatin with BSA (BSA Control) was at 88.88 ±2.24% and amphotericin B-nystatin with protein elution buffer (Elution Control) was at 83.23 ±2.53%. Other components of the fungicides, such as the deoxycholate present in the amphotericin B solution, were also tested and found to have no impact on survivability at the levels used in these assays (Figure S11). Finally, all three treatments with recombinant splice variants (which includes amphotericin B and nystatin) displayed decreased survivability (albeit at different levels between 6–14%) relative to control treatments (pre-treated with At1g74130 (L) displayed 13.93 ±1.16% survival, (S) at 6.92 ±4.39%, and (M) at 10.08 ±2.53%). Yeast cells pre-treated with recombinant At1g74130 variant proteins and then treated with both amphotericin B-nystatin exhibited the smallest colony sizes (Figure 9B). Only pre-treatments with recombinant splice variants and amphotericin B resulted in higher sensitivity to nystatin. Protein
Figure 9. Activity assays using recombinant At1g74130 splice variant proteins as exogenous additives and yeast cells. (A) The typical treatment matrix used in this study is shown. The check marks in the columns indicate the presence of a treatment component. The variant protein tested in Figure 7 is labelled only as L, S, and M in the matrix. The final treatment volume in all cases was 100 µL. (B) The corresponding plate is shown for each of the treatments depicted in panel A (labelled according to treatment numbers 1 to 9). (C) Cells (resulting colonies) surviving the different treatments are depicted as Percent Survival in the graph. The treatments are in the same order as presented in panels A and B. The error bars represent standard deviations (minimally n=3).

Pre-treatments without the delivery agent amphotericin B behaved like the controls (Figure S11).

Changes to ampicillin sensitivity in bacteria were also observed using the transient approach. As a commonly used delivery component in many drug applications, DMSO was utilized here as the protein delivery agent in place of amphotericin B. Bacteria were pre-treated with DMSO and recombinant proteins before testing ampicillin sensitivity. Relative to the untreated (media only) and mock treatment (all components without recombinant proteins), pre-treatments with exogenously added At1g74130 splice variants decreased the number of colonies (a proxy for cells surviving the treatment) at the time of plate growth documentation (Figure 8D and Dataset 2). The mock treatment did not differ significantly (T-test, P = 0.36) from the no treatment control, indicating that the components used in the buffer did not
contribute significantly to the enhanced level of ampicillin sensitivity. Relative to the mock treatment or no treatment control, the pre-treatment of bacteria with recombinant protein additives At1g74130 (L), (M), or (S), exhibited significant reductions in colony numbers (T-test: P = 0.022, P = 0.026, P = 0.029, respectively). The No Treatment control using 1.25 mg/ml ampicillin without DMSO, represents the reference point of 100% survival. The Mock Treatment without protein additives and with DMSO resulted in 95.85 ± 6.08% survival. The treatments (5% DMSO and 1.25 mg/ml ampicillin) and pre-treated with At1g74130 (L), (M), or (S) resulted in 71.36 ± 8.96%, 79.54 ± 3.10%, and 77.00 ± 1.27% survival, respectively.

The functionality assays used for At1g74130 were applied to splice variants from two other categories of rhomboid proteins. One splice variant pair was derived from Arabidopsis At1g25290 (named (L) and (S)) and another variant originated from human Ubac2. At1g25290 was from the “Active Rhomboid Proteases” category and Ubac2 is from the “Other Inactive Rhomboid Proteins” category. The overall results for variants from these two other categories indicate functionality as well in our assay settings (select assay results are reported here).

For the At1g25290 splice variants (L) and (S), similar responses were observed in the exogenous additive-transient setting, albeit with differences in impact from that observed for the At1g74130 variants. Functionality was displayed in both bacterial (Figure 10A and Dataset 4) or yeast cell settings (Figure 10B and Dataset 4). The outcomes between the two At1g25290 splice variants were themselves different. The different sensitivity levels displayed by (S) relative to (L) suggest that the phenomenon observed is attributed to the added recombinant rhomboid variant (that their differences were derived from alternative splicing) and not to other components in the mixtures.

The human Ubac2 splice variant tested is a fusion between a rhomboid protein sequence (considered a pseudoprotease) and ubiquitin-associating domains. Like the above phenomenon, recombinant Ubac2 variant proteins showed functionality as an additive in bacterial and yeast assays, albeit at a more modest level of influence on antimicrobial sensitivity (Figure 11; Dataset 5).

In whole, not all variants will show functionality in the different test settings. The different assays does however provide evidence that splice variants from different rhomboid categories possess functionality, supporting the notion derived from the comparative analysis, that alternative splicing provides a mechanism for diversifying the numbers of working rhomboid proteins, and the roles of rhomboid proteins play in a particular system.

![Activity assays using At1g25290 splice variant proteins as exogenous additives to bacteria and yeast cells.](image)

**Figure 10.** Activity assays using At1g25290 splice variant proteins as exogenous additives to bacteria and yeast cells. (A) Recombinant At1g25290 splice variant proteins L and S were tested for activity (for enhanced sensitivity to ampicillin in this case) as exogenous additives and bacteria. The assays were conducted in the same manner as that shown in Figure 8 (from untreated to mock to added proteins). Cells (resulting colonies) surviving the different treatments were then assessed and represented as Percent Survival. The bar graphs are arranged to the corresponding representative results, the resulting agar plates. The error bars represent standard deviations (n=4, T-test, P=0.01). (B) Recombinant At1g25290 splice variant proteins L and S were also tested for activity (for enhanced sensitivity to nystatin in this case) as exogenous additives and yeast. The assays were conducted in the same manner as that shown in Figure 9 (from untreated to mock to added proteins). The organization of panel B is the same as in panel A, except for yeast and nystatin. The error bars represent standard deviations (n=4, T-test, P=0.01).
The error bars represent standard deviations (n=4, T-test, P=0.1). The assays were conducted in the same manner as that shown in Figure 8 (from untreated to mock to added proteins). Cells (resulting colonies) surviving the different treatments were then assessed and represented as Percent Survival. The key results comparing the splice variants At1g25290 (L) and Ubac2 are shown in this panel. The error bars represent standard deviations (n=4, T-test, P=0.1). (B) Recombinant At1g25290 splice variant protein L and Ubac2 were also tested for activity (for enhanced sensitivity to nystatin in this panel) as exogenous additives and yeast. The assays were conducted in the same manner as that shown in Figure 9 (from untreated to mock to added proteins). The organization of panel B of this figure is the same as in its panel A, except for yeast and nystatin. The error bars represent standard deviations (n=4, T-test, P=0.1).

***Discussion and conclusions***

Alternative splicing is used by many organisms to control and to diversify protein function. Historical examples include human tropomyosin, human kallikreins (secreted serine proteases), and fungal Ski7/Hbs proteins[23,24]. The same appears to be possibly happening with rhomboid genes, but despite witnessing alternative splicing as a mechanism for diversifying functionality in *Arabidopsis* and human breast cancer cells[18,22,24,29,45], the number of demonstrated cases remains limited. We were thus interested in assessing how often this happens using the information available in the different genetic databases. Our comparative survey of current databases was focused specifically on functionality, as opposed to function, to capture slight changes (potential changes at this juncture) to functionality, such as those reported above[23,24]. We also wanted to determine the extent of alternative splicing within the different rhomboid gene systems of a particular eukaryotic organism as well as between different species. We thus limited our analysis only to splice variant entries in current RNA sequence databases of six eukaryotic organisms, ones used widely as experimental models. It was important to limit our analysis so that we can address the issue of extent and then assess how alternative splicing could be used to modify the functionality of distinct regions of the affected rhomboid proteins of our data set, especially regions with defined purposes. We then assessed the overall notion of the comparative findings by testing a selection of variant proteins from three different categories.

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**Figure 11.** Activity assays using a select Ubac2 splice variant protein as exogenous additives to bacteria and yeast cells. (A) Recombinant splice variant proteins At1g25290 (L) and one Ubac2 were tested for activity (for enhanced sensitivity to ampicillin in this panel) as exogenous additives and bacteria. The assays were conducted in the same manner as that shown in Figure 8 (from untreated to mock to added proteins). Cells (resulting colonies) surviving the different treatments were then assessed and represented as Percent Survival. The same appears to be positively happening with rhomboid genes, but despite witnessing alternative splicing as a mechanism for diversifying functionality in *Arabidopsis* and human breast cancer cells[18,22,24,29,45], the number of demonstrated cases remains limited. We were thus interested in assessing how often this happens using the information available in the different genetic databases. Our comparative survey of current databases was focused specifically on functionality, as opposed to function, to capture slight changes (potential changes at this juncture) to functionality, such as those reported above[23,24]. We also wanted to determine the extent of alternative splicing within the different rhomboid gene systems of a particular eukaryotic organism as well as between different species. We thus limited our analysis only to splice variant entries in current RNA sequence databases of six eukaryotic organisms, ones used widely as experimental models. It was important to limit our analysis so that we can address the issue of extent and then assess how alternative splicing could be used to modify the functionality of distinct regions of the affected rhomboid proteins of our data set, especially regions with defined purposes. We then assessed the overall notion of the comparative findings by testing a selection of variant proteins from three different categories.
The overall evidence from the comparative analysis supports the hypothesis that alternative splicing could be used to diversify rhomboid functionality in a number of cases. This is especially the situation in human, mouse, and Arabidopsis, organisms with relatively high numbers of rhomboid or rhomboid-like genes. Currently, there is a total of 95 entries for 13 human rhomboid genes that reflect alternative splice products, 53 for 13 mouse genes and 40 for 22 Arabidopsis genes. These splice variants were also not limited to a particular rhomboid category. The diversification appears to occur generally in distinct groupings across the entire rhomboid protein sequence. Although the comparative data suggest potentially impactful changes to functionality, this speculation remains limited to linear protein sequences and motifs. Therefore, we next tested the possible changes to functionality using currently available predictive tools for 3-D protein structures, despite the highly speculative nature of these tools. This analysis was focused strictly on how changes could theoretically impact such a rhomboid structure. Because these assessments are judged as being too speculative, the outcomes of these tests are provided only as Supplementary Material (Supplementary File 1 and Figure S1–Figure S8). These outcomes may be useful for guiding future structural studies as well as validating the outcomes through extensive experimentation. The notion of diversification in functionality by alternative splicing mechanisms was tested experimentally using recombinant proteins of six different splice rhomboid variants from three different categories, Active Rhomboid Proteases, Inactive Rhomboid Proteins, and Other Inactive Rhomboid Proteins. These splice variants represent different structural changes from active sites, to truncations, to fusions.

Based on the compiled comparative data, some of the potential impacts were quite extensive and obvious. Some of the more obvious ones appear to arise from subtle changes to the protein sequence, such as the introduction or removal of a few residues. Many impacted important structural motifs, sometimes from afar or indirectly (Figure 2–Figure 6 and Figure S1–Figure S8). Although the degree of amino acid sequence conservation of rhomboid and rhomboid-like proteins is relatively low between species and types, there are distinct conserved residues/motifs that serve the same important functions. Some of the functional motifs potentially impacted did include motifs of known functions like the L1 loop, the TMD5-L5 cap, and the catalytic dyad region. Such situations are likely to bear significant consequences with respect to functionality. For instance, the L1 loop has been the focus of several studies because it contains one of the conserved motifs outside of the catalytic cluster. Although the function of the L1 loop is not entirely understood, the importance of the loop on protease functionality was demonstrated by mutagenesis experiments. Normally, the L1 loop is partially embedded in the membrane. Mutation of the conserved WR motif in the L1 loop of GlpG decreased proteolytic activity, suggesting a modulatory role for the loop. Additional evidence suggests a regulatory role for the L1 loop, and this role is linked to the formation of a rigid L1 loop structure. If the L1 loop serves as an anchor to the lipid bilayer, alterations to this structure could modulate the protein’s enzymatic activity with substrates. In 2007, Baker and coworkers found an enhancement of proteolytic activity with their set of mutagenized L1 loop experiments, which further demonstrates the link between L1 loop and functionality. Such outcomes are certainly possible and were observed in the 3-D models predicted for the different splice variants tested here (Figure S7).

Similar outcomes were also observed for splice variants involving changes to the L5 cap and TMD5 region (Figure S8). In addition to experiments aimed at the L1 loop, Baker and coworkers also carried mutagenesis-based experiments on the TMD5 region and observed enhancement of cleavage activity with some of the structural changes in TMD5. It was hypothesized that destabilization of the TMD5 helix allowed enhanced substrate entry. TMD5’s destabilization is believed to alter its configuration by changing its angle/tilt and proximity to neighboring helices. This in turn alters the efficiency of gating by this region. Alterations to the efficiency of gating then in turn affects proteolytic activity. This is because, normally, when the TMD5 is positioned in the ‘open’ conformation, the TMD5 helix pulls the L5 loop outward. The movement of the L5 cap structure allows substrate entry into the catalytic cavity. The open conformation of the L5 cap is also believed to permit the entry of water into the catalytic cavity. Therefore, alterations to TMD5 and the L5 cap could potentially change features that affect substrate entry. The predicted outcomes in the examples shown in Figure S8 could possibly manifest in a similar manner with equally consequential effects. The structural outcomes revealed in our theoretical assessment of splice variants are thus likely to represent a mechanism for diversifying rhomboid functionality, since these alternative splice variants likely exist in the organisms studied.

Based on a combination of the previous findings for the two Arabidopsis plastid rhomboids, the human RHBD2, and the trends revealed in this study, the overall evidence suggests that alternative splicing is a functionally significant mechanism for diversifying rhomboid functionality. This means that splice variants of rhomboids and rhomboid-like proteins likely exist simultaneously in the cell or sub-cellular compartment. Like the two Arabidopsis plastid proteins, the alternative splice variants are likely co-expressed, modulated relative to each other to respond to the cell’s needs, and interacting in some manner. The possibility of interactions between rhomboid units themselves has been reported by Wu et al. for the bacterial rhomboid protease GlpP. Such interactions with different rhomboid variants/forms and populations could therefore manifest in a number of ways that affect rhomboid functionality. The theoretical approach used in this study, and the predicted outcomes that may arise are thus not without merit, and should be considered as guidance for further experimentation. The possibilities, such as the ones discussed in the above examples, are observed experimentally in other studies and in functionality assays conducted for our select splice rhomboid variants. There are a number of other experimentally tested examples where truncations have been observed to impact functionality. In addition to rhomboid proteins, examples of other types of proteins include those recently reported by Stoddart et al. for an integral membrane pore, and by Quemeneur et al. where shape influences protein mobility within membranes. Whatever the situation may be for rhomboids, it is clear that it is necessary to characterize splice variants for each rhomboid and to determine how splicing influences rhomboid functionality. This would be
important for elucidating how the different rhomboids work as a network.

Data availability

Dataset 1: Raw data underlying the results in Figure 7. Figure 7B. Disk Assay Area Measurements of Growth (Sensitivity) in the Presence of Nystatin for Transgenic Yeast Lines Expressing At1g74130 Splice Variants; Figure 7C. Percent Survival (Cell/Colony Counts) of Various Transgenic Yeast Lines When Treated with Nystatin and Amphotericin B; Figure 7D. Densitometry Assessment of Immunoblots Developed for Assessing Mgm1 Levels and Ratios in Mitochondria Isolated from Transgenic Yeast Lines Expressing At1g74130 Splice Variants. 10.5256/f1000research.13383.d204676

Dataset 2: Raw data underlying the results in Figure 8. Figure 8A–B. Cell/Colony Counts of Various Transgenic Bacterial Lines When Grown on Plates with Increasing Ampicillin Concentrations; Figure 8D. Assays to Determine Percent Survival (Cell/Colony Counts) of Bacterial Cells When Treated Exogenously with Recombinant At1g74130 Splice Variant Proteins. 10.5256/f1000research.13383.d204677

Dataset 3: Raw data underlying the results in Figure 9. Figure 9C. Assays to Determine Percent Survival (Cell/Colony Counts) of Yeast Cells When Treated Exogenously with Recombinant At1g74130 Splice Variant Proteins. 10.5256/f1000research.13383.d204678

Dataset 4: Raw data underlying the results in Figure 10. Figure 10A. Assays to Determine Percent Survival (Cell/Colony Counts) of Bacterial Cells When Treated Exogenously with Recombinant At1g25290 Splice Variant Proteins; Figure 10B. Assays to Determine Percent Survival (Cell/Colony Counts) of Yeast Cells When Treated Exogenously with Recombinant At1g25290 Splice Variant Proteins. 10.5256/f1000research.13383.d204680

Dataset 5: Raw data underlying the results in Figure 11. Figure 11A. Assays to Determine Percent Survival (Cell/Colony Counts) of Bacterial Cells When Treated Exogenously with a Recombinant Ubac2 Splice Variant Protein; Figure 11B. Assays to Determine Percent Survival (Cell/Colony Counts) of Yeast Cells When Treated Exogenously with a Recombinant Ubac2 Splice Variant Protein. 10.5256/f1000research.13383.d204681

Competing interests

Parts of this study may be considered as being potentially related to material contained in a patent application (US 2016/0129078 A1). There are no other competing interests disclosed at the time of submission.

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Supplementary Material

Supplementary File 1: Supplementary notes for Figure S1 to Figure S8; References cited in the supplementary notes for Figure S1 to Figure S8; Supplementary figure legends for Figure S1 to Figure S11.

Click here to access the data.

Supplementary Tables S1–10

Click here to access the data.

Figure S1. Examples of alternative splicing and their potential impacts on the structure of the amino terminal region.

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Figure S2. Examples of alternative splicing and their potential impacts on the structure of the carboxyl terminal region.

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Figure S3. The potential impact of alternative splicing on the structure of the Arabidopsis rhomboid At3g17611.

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Figure S4. The potential impact of alternative splicing on the structure of the Arabidopsis rhomboid At3g53780.
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Figure S5. Examples of alternative splicing and their potential impacts on the L1 loop structure.
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Figure S6. Examples of alternative splicing and their potential impacts on the L5 Cap-TMD5 structure.
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Figure S7. The potential impact of alternative splicing on the structure of the Arabidopsis plastid rhomboid At1g25290.
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Figure S8. Examples of alternative splicing and their potential impacts on the structure of the catalytic region.
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Figure S9. Immunoblot images used for analyzing the impact of different rhomboid combinations on the Mgm1 ratios presented in Figure 7D.
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Figure S10. Immunoblot images used for analyzing the impact of rhomboid variants on β-lactamase production and secretion presented in Figure 9C.
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Figure S11. Assay testing the combination of “Inactive” plastid rhomboid protein variants and deoxycholate in yeast without amphotericin B.
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References


18. Zou H, Thomas SM, Yan ZW, et al.: Human rhomboid family-1 gene RHBD1 participates in GPCR-mediated transactivation of EGFR growth signals in...


The authors Powles and Ko now provide a revised version. Although this version is improved compared to the previous version, the authors did not follow my suggestion to revise the manuscript except for the sentences, which I explicitly stated. The manuscript is still full of awkward phrases, where it is clear what the authors want to say but which read wrong as they are written. It is not my but the authors' task to carefully go through the manuscript and correct all these issues. Citations are still extremely unbalanced. In the introduction and discussion, citations are pooled, while at many statements that would warrant citations those are missing. I also cannot accept the sequence-based analysis as it is. If the authors prefer not to revise it I would suggest to completely remove it or boil it down to a single schematic figure and at most half a page of text. As it is, it is completely useless for further research: researchers not aware of the problems with sequence analyses will be misguided, and researchers interested in sequence aspects will need to do a correct analysis anyway. The supplementary Figures S1 to S8 do not show any useful information (the figure legends do not describe what is shown, the structures in “A” do not correspond to the sequences shown in “B”) and need to be completely removed. The authors do not understand what RNA-seq mapping means and everything in the manuscript mentioning RNA-seq needs to be removed. The authors write “Entries were retrieved from databases for cDNA/EST sequences, RNA-seq,…”, but where are these entries? Where is the list of reads from SRA archives supporting any of the alternative splice variants? Which cDNA/EST clones support which splice variant? Many (especially the human) splice variants result from COMBINATIONS of alternatively spliced exons, e.g. from 2 alternatively spliced exons you might get 4 different transcripts and this exponentially increases with the number of alternatively spliced exons per gene. Where is the evidence (full-length cDNA) that all these combinations are found? Are there coupled splicing events? How does the number of “entries” reported (e.g. 95 for human) correspond to the number of possible combinations, and by which evidence are certain combinations excluded? By comparing human and mouse, isn’t it just a database problem that mouse contains less entries than human, although the total number of potential combinations is identical?

In their response letter, the authors write “We did not include references dealing with alternative splicing because the study was not intended to be at this level, but working at the level of surveying and comparing available entries and what these entries may reflect at the protein level. As described above, substantial revisions were focused on clarifying the intent of the work to avoid this impression. This reasoning applies to why gene structures and splicing events were not provided.”. If the authors do not want to deal with alternative splicing at the gene structure level, than the provided data/figures are even more misleading and misleading. What is the benefit for the reader to have all these figures (main figs and supplementary data/figs) if these are not based on and supported by real data? Either, do a thorough
analysis based on gene structures and real RNA-seq mapping, or remove all the speculations on predicted sequences and structures (text, figs, supplements). Do the authors know which software was used to generate these predictions? Re-run the gene prediction software and you will get more predictions to speculate on.

=> Abstract, Results section, first sentence: “A comparison of database entries of six widely used eukaryotic experimental models […] revealed robust usage of alternative splicing…”. I think that the word “reveal” is overstating the findings, the database entries just “suggest” that alternative splicing might happen. What is “robust usage of alternative splicing”?

=> Introduction, first paragraph: Lemberg and Freeman do not mention “Derlin” in their paper at all. The current phrasing of this sentence is misleading in this respect. As far as I understand the Lemberg and Freeman paper (see Fig. 4 there), they distinguish 3 types of rhomboids, and the proteolytically active ones further subdivide into two subgroups. The sentence by Powles and Ko “Phylogenetic studies, such as that conducted by Lemberg and Freeman (2007), suggest that rhomboid genes can be divided into two subgroups encoding proteolytically active secretases- and Presenilin Associated Rhomboid Like (PARL)-type rhomboid forms, and two subgroups of catalytically inactive forms (“non-proteases”) such as iRhoms, Derlins, and other distantly related forms." reads as if there were 2 main groups (proteolytically active and catalytically inactive) and that the catalytically inactive further subdivide into to two subgroups (mentioning even three subgroups, iRhoms, Derlins and other distantly related forms). This is not a clear description how this protein family is organized and which groups/subgroups are distinguished. I haven’t read the other 4 cited papers, just the one by Lemberg and Freeman, thus I cannot suggest how to rephrase this sentence to be consistent with the literature. In the end, it should be clearly stated which and how much major subgroups are distinguished, and which of these are further distinguished in minor subgroups, and which of the classifications is based on which publication. The next sections are at least not in-line with the Lemberg and Freeman classifications, as they state “The active category…” implying no subgroups and “The inactive categories…” implying multiple subgroups.

=> The citation of previous literature in the Introduction section is still not adequate. While the citations in the section about “active rhomboids” is appropriate putting citations to the listed functions, in the section about “inactive rhomboids” all citations are just combined at two occasions, although many more different functions of rhomboids are listed than in the “active” section. It should be the authors’ and not the readers’ task to put citations at appropriate statements. As it is, there are multiple citations for very general statements, while all the detailed information is not supported by citations.

=> “…At1g741301, also exists as three splice variants…”: What does “also” mean here? Where is the reference to other rhomboids with 3 splice variants?

=> “…changing from a seven predicted to six transmembrane structure”: I don’t understand that phrase, this does not make any sense.

=> “Functionality differences…”: What does this mean? There are many more awkward phrasings throughout the manuscript, that I will not list here. I highly recommend the authors to go through the manuscript again and improve the phrasing. There are examples in almost every second sentence… (e.g. “A limited selection of alternatively spliced variants were…”, “In many cases, alternative splicing events tend to influence one motif, but there were a number of cases where the impact may affect one or more motifs…”, etc. etc.)

=> “…we conducted a comparative analysis of entries available in the genetic databases…”: Did you
really look in “genetic” databases (and if, I would be very curious to know which exactly) or genomic databases?

=> Methods section: “For instance, a frequent occurrence is an alternative splicing event which occurs in a particular motif that would likely impact the adjacent linker region.” Does this sentence provide any useful information? Is it related to any “method”? Does it implicate that “particular motifs” as part of alternative splicing events are only located in domains (e.g. TMs) and not in linker regions? Does it implicate that these motifs never impact the motifs themselves or the corresponding domains but only the linker regions? Wouldn’t such events rather impact spatially close regions (e.g. tertiary structure) instead of regions connected by sequence succession? I highly recommended in my first review to revise the Methods section with respect to removing unnecessary information not related to any Method, and instead provide a description what was actually done with which software (this includes mentioning all the software versions! BLAST v.1.0 is very different from BLAST v.2.28, for example). The Methods section needs further shortening (remove all what is not a method) and considerable rephrasing (to make exact statements; e.g. what is a “Linear protein alignments…”? Are there circular or other alignments? What are “Structural predictions…”, “Three-dimensional predictions…”?).

=> Shouldn’t it be “selected” instead of “select” in the title and at many other occasions in the manuscript, e.g. “Select proteins were synthesized…”?

=> I only looked at Figure S1 in detail, and because this is already that strange and misleading I stopped and did not look at the other supplementary figures. From its title “Examples of alternative splicing and their potential impacts on the structure of the amino terminal region” the reader would suspect to see a structure of the N-terminal region, and structures where this region is changed because of alternative splicing events. At first glance, Figure S1 shows 13 structures in “A” and an alignment of 13 sequences in “B”. However, the structure denotes TMD3-6, which are not part of the N-terminal region and not part of the red-boxed region of the alignment, which shows the differences due to alternative splicing. Where is the N-terminal region in the structure? According to the figure legend, “Key residues of the catalytic site are noted by colorized letters that match the colorized regions of the generated 3-D models”, but in the 3D-models all secondary structural elements are coloured, not the key residues. There is no way to identify these “key residues” anywhere in the structure. According to the alignment shown in “B” there are three splice events: i) there could be a short elongation at the N-terminus, ii) there is a differentially included region in the middle, and iii) almost the entire N-terminal region can be missing. But where do I see this in the structures? Sequences x1, x2, and x6 should have identical N-terminal sequences, but the corresponding structures in “A” are different. Or do numbers in “A” not correspond to numbers in “B” as stated in the figure legend? Same for sequences x3, x4, and x5, which are identical but do show different structures. Why do have the sequences x10, x11, and x12, which almost completely miss the N-terminus, structures at all? The sequences for the N-termini of all 13 variants are identical (except for including/excluding certain regions), why are the structures for these N-termini different?

**Competing Interests:** No competing interests were disclosed.

**Referee Expertise:** bioinformatics, gene prediction, eukaryotic genomes, alternative splicing, structural biology, protein expression, biochemistry

I have read this submission. I believe that I have an appropriate level of expertise to state that I do not consider it to be of an acceptable scientific standard, for reasons outlined above.
The manuscript presents original materials on alternative splicing of rhomboid proteins. The authors discuss potential impacts on the rhomboid protein structure. The paper has good illustration and rich Supplementary materials. A lot of work done. Thus, it is technically sound.

But the presentation has to be improved. A lot of unnecessary literature citations (such as 2-31). Thus, the work does not have clear literature citations.

Choice of model objects is not clear. Alternative splicing prediction depends on computer tools. If we compare splice variants of the same protein (homolog) from different species, the parameters of prediction should be comparable. Some model organisms, such as human and mouse, have much more experiments and works, and just better gene annotation than for plant genomes. Thus, we can't directly compare number of splice variants. Thus, I consider the statistical analysis and its interpretation only partly appropriate. And not fully reproducible - to answer the question ‘Are all the source data underlying the results available to ensure full reproducibility?’

‘Periodic comparison’ or ‘multi-year analysis’ of the information on the proteins family databases should be explained. What is the time for database releases? What kind of trend in functional potential could annotation of these proteins prove? I recommend remove that part of the paper.

The study is based on the work by the authors in 2012 on the Arabidopsis thaliana homologs containing alternative splice variants. The current article should show novel materials and conclusion. Comparison by the available information updates on these homologs from model organism databases is a weak idea. If you wait longer, more model species will be functionally annotated, and definitely some new splicing variants found. So, it could be either predictive model, or just description of all data available now on rhomboid proteins.

‘Multi-year analysis’ is a redundant term - I recommend remove such part of text, or reformulate it. Overall, the article presents novel ideas on the cross-species comparison of alternative splicing. This science area is growing and we might expect new evidence for alternative splicing from genome sequencing projects.

The conclusions drawn by the authors are adequately supported.

Thus, this work is of interest for F1000Research readers. Thus, I recommend ‘Approved with Reservations’ status.

Is the work clearly and accurately presented and does it cite the current literature? Partly
Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Partly

If applicable, is the statistical analysis and its interpretation appropriate?
Partly

Are all the source data underlying the results available to ensure full reproducibility?
Partly

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Referee Expertise:** bioinformatics, genomics

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 04 Apr 2018

**Kenton Ko, Queen's University, Kingston, Canada**

Thank you for taking the time and effort to review our submission. This is very much appreciated!

The feedback provided from the different reports is valuable to us and will guide our revisions for the next version of this contribution.

We will be in touch soon with the next version.

**Competing Interests:** None

Author Response 22 May 2018

**Kenton Ko, Queen's University, Kingston, Canada**

Dear Dr. Orlov,

Thank you for agreeing to review version 1 of our submission and helping us with your assessment. This is much appreciated!

We now have version 2 available for your further assessment. Based on the feedback, Version 2 was revised substantially to address the aspects raised and to enhance clarity in the areas of writing style and additional details.
As per your suggestions, improvements were made to the presentation by removing references that may not be necessary to the readers. Removal of these extra references allowed us to revise the Abstract, Introduction, and Methodology sections to enhance clarity with respect to the study’s focus or purpose. The title was also modified slightly to reflect these revisions.

The choice of model organisms used was explained better by revising the style of writing/presentation and by characterizing the work as a comparative analysis or survey of available entries, as opposed to a direct comparison of splice variant numbers between species and its interpretation in this way. We hope that we have interpreted this suggestion correctly. This aspect, we believe, was also raised by Reviewer 1, Dr. Kollmar.

We have revised the Methods section substantially to clarify the approaches used and to provide more needed details concerning the analysis of available database entries and the functionality assays using recombinant proteins. Revisions were also made to the Source Datasets and the Supplementary Tables to enhance their presentation, to provide more descriptive titles and related aspects. We hope that there is now sufficient methodological details for others to follow.

The multi-year facet of the database work was removed to avoid confusion and to enhance clarity. This removal allowed us to streamline the database analysis protocols and provide more details of the tools used. The Results and Discussion sections were modified similarly to reflect these changes. The removal of the multi-year aspect in combination with the above revisions should help the readers interpret the work more clearly.

Sincerely,

Kenton Ko and Josh Powles

Competing Interests: No competing interests were disclosed.

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Lu-Yuan Li

State Key Laboratory of Medicinal Chemical Biology, College of Pharmacy, Tianjin Key Laboratory of Molecular Drug Research, Nankai University, Tianjin, China

The rhomboids are a very large family of genes found across all kingdoms from bacteria to plants to animals. Based on what we know about the functions of these genes, which unfortunately is not very much at the moment, they often play critical roles in life wherever they are involved. Also interesting is that their functions are highly diversified, as mentioned in this paper. For instance, the human rhomboid family 1 gene, RHBDF1, was found to be expressed at significantly higher levels in breast cancer tissues than in normal breast tissues, with functions as different as promotion of G-protein coupled receptor mediated transactivation of epidermal growth factor receptor (reference 1), or protection of hypoxia-inducible factor 1-alpha from degradation under hypoxic conditions (reference 2); both functions are consistent with
facilitating tumor growth, however. RHBDF1 belongs to the group of so-called inactive rhomboids (iRoms) because of an apparent lack of protease activities associated with otherwise enzymatically active rhomboids. This group of rhomboids, having lost their abilities to catalyze proteolysis reactions due to mutations, seem to have destined in evolution to perform other functions, such as those of chaperones, with their abilities to bind to a variety of proteins conserved and utilized in assisting protein folding, transportation, and degradation.

The paper by Powles and Ko is interesting as it addresses a fundamental cause of the massive variations of the functions of the rhomboid gene products. The authors carried out an extensive data mining operation to reveal complex differential splicing patterns of the rhomboid genes. Their findings indicate that differential splicing is common and substantial within one gene as well as throughout the gene family. Apart from the multiple transmembrane domains, the rhomboid proteins often possess a large N-terminal domain before the first transmembrane domain and a sizable loop between the first and second ones. The N-terminal and the loop are likely located on opposite side of the endoplasmic reticulum, Golgi, or plasma membrane of the cell, indicating that they have opportunities to interact with different biomolecules. In addition, in the case of RHBDF1 there are a number of amino acid residues within the protein molecule that could be subjected to post-translational modifications such as phosphorylation and glycosylation. It would therefore seem highly likely that frequent and extensive splicing of the gene transcripts exerts considerable impact on the protein structures and thus their functions. Changes in the gene transcripts could also bring about alterations in terms of modulations by micro RNA and other non-translational means. The effort by the authors is very useful to put together a data base of immense and complex differential splicing patterns of nearly the entire rhomboid gene family. The findings should be beneficial to researchers in this field, even though many of the conclusions are speculative, as pointed out by the authors, because of the lack of our knowledge on the structures and functions of most of the individual products of this enormous gene family.

References

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes
Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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**Author Response 04 Apr 2018**

**Kenton Ko, Queen's University, Kingston, Canada**

Thank you for taking the time and effort to review our submission. This is very much appreciated!

The feedback provided from the different reports is valuable to us and will guide our revisions for the next version of this contribution.

We will be in touch soon with the next version.

**Competing Interests:** None

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**Author Response 22 May 2018**

**Kenton Ko, Queen's University, Kingston, Canada**

Dear Dr. Li,

Thank you for agreeing to review version 1 of our submission and for your assessment. This is much appreciated! We now have version 2 available for viewing. Based on the feedback, Version 2 was revised substantially to address the aspects raised and to enhance clarity for the readers in the areas of writing style and additional details. The nature of the work reported was not altered by the revisions.

The presentation of the study’s purpose and approach used was revised for clarity. This was achieved first by removing most of the references listed from 2 to 26 so that the references cited were more obviously directed at the study, as opposed to providing general background of the wider field. Accompanying revisions were then made to the Abstract and Introduction.

The Methods section was revised substantially to clarify the approaches used and to provide more needed details concerning the comparative analysis of database entries and the functionality assays using recombinant proteins. The title was also modified slightly to align with these revisions. The Results and Discussion sections were modified similarly to reflect the changes. The multi-year facet of the database analysis was deemed unnecessary and removed. Revisions were also made to the Source Datasets and the Supplementary Tables to enhance their presentation. Collectively, these revisions should help the readers interpret the work more clearly with the additional details and streamlining of the protocol for the database work.

Sincerely,
Joshua Powles and Kenton Ko conducted a study on rhomboid serine proteases with respect to their alternative splice variants. The study is based on a discovery of the same researchers in 2012 that two of the Arabidopsis thaliana homologs contain alternative splice variants. The manuscript displays several conceptual problems that would demand a major revision if touched. Therefore I suggest the authors to completely revise all in silico analyses and completely re-write the paper. The problems are as follows:

=> The authors provide much unneeded information for certain aspects (which would just be a style problem) but, most importantly, miss a lot of essential information to recapitulate and redo their analyses. Example: The authors state in the Methods section: “All variant RNA sequences were assumed to result in the generation of functional proteins, including extensively truncated products.” This, however, is not a method. A method would be that the authors write, that they used tool-x or tool-y to translate the RNA sequences, which they subsequently analysed by tool-z. Instead, the authors continue: “The resulting predictions did not indicate that this assumption would not hold true.” What does “resulting” mean here? Which tool was used? A translation of an RNA is usually not termed “prediction”. Which tool was used for the verification? Apart from these style problems, the authors did not provide any evidence that the translations “do” result in functional proteins. It still is just their assumption. In terms of missing information to recapitulate/redo the analyses, the authors do not provide any information on: A) how the set of starting proteins were assembled (search by name? Using BLAST?), B) how the entries were assessed for alternative splicing (on RNA? On protein level? By alignment?), C) the authors claim that their analyses required comparisons between genomic sequences but do not provide any genomic sequence in the main text or supplements, D) the authors claim “alignments with current models of rhomboid proteins” without providing the tool used for doing these alignments nor providing a list or reference for the “current models of rhomboid proteins”, E) the authors state that they used Ni-NTA chromatography for protein purification without providing any buffer. While column washing-buffers likely do not have any effect, the elution buffer has. Was the elution done in a gradient? By a step-wise process? Just by eluting with 500 mM Imidazole? Which pH? Was the purified protein dialysed afterwards? This is essential information, because the proteins were subsequently used in other assays, and these might be compromised by pH or imidazole concentration. F) the authors do not provide any information on ampicillin-concentration used for selection, nor any cell lysis procedure. G) was the purified protein always used directly? Frozen and reused later? How long is the protein stable in which buffer (pH? Imidazole?)? H) I do not understand this sentence: “If used, the rabbit polyclonal anti-rhomboid protein (At1g74130) antibodies were established by our lab and validated as reported in Powles et al. 51.” Did the authors use this antibody for the western blots? Why did they “validate” the antibody again? I) which media were used for yeast cell growth? Just writing “Cells were grown in glucose-supplemented media” does not make the experiment reproducible. Which glucose concentration? What is a “complete medium”? J) The authors experimentally analyse
three variants for the At1g74130 gene, termed L, M, and S, but it is not described anywhere to which splicing events these transcripts belong. Every study that I know that deals with alternative splicing contains a gene structure scheme and shows which exons would be present in which transcript. This is just an incomplete list of examples, where absolutely essential information is missing. The authors would need to check every sentence in the Methods section to see, whether it is really describing a method, and they need to check where and which information is missing.

=> I have looked at the provided accession numbers for the genes/proteins. The numbers for human and mouse all represent "predicted transcripts", thus these could all be wrong. I recommend the authors to read a few papers about gene prediction software and pipelines, they will notice that on average every prediction in e.g. human, Drosophila, C.elegans contains 1 wrong exon (e.g. the paper from the eGASP comparison of gene prediction software on these 3 species would be a good starter). Prediction of alternative splice variants is even more error-prone. Thus, the authors should only use transcripts with cDNA evidence in their study (and keep in mind, that cDNA/EST data also sometimes contains errors from e.g. missplicing). Discussing just gene predictions is complete artificial and fiction. The authors could, for example, download a few RNA-seq datasets from the ENCODE and modENCODE data and do the RNA-seq mapping themselves to validate the gene predictions. If the authors cannot do this, they should only analyse and discuss transcript variants for which they find cDNA/EST evidence (and of course the accession numbers for these cDNA sequences need to be given). If the authors did a RNA-seq mapping themselves they could also provide some measure for the likeliness that a suggested variant is a true variant or the result of missplicing, transcription errors, etc. E.g. if thousand of reads are found for a certain gene, is it likely that a variant is a true variant and functional if only supported by 1-2 reads? Or are these 1-2 reads rather representing missplicing and other errors?

=> The authors claimed several times that they re-did the analysis every 6 months since 2012. Wouldn’t it be a much better way to just look first whether updates on these species were made available at all before spending time in redoing an analysis? I know that these updates only happen occasionally, and not even every second year. Also, the techniques completely changed since about 2010. At least I am not aware of any major study providing new EST/cDNA datasets for the species studied here. All the new data since about 2008 is generated as RNA-seq data. Thus, which further cDNA evidence did the authors expect for the transcripts since 2012 so that they decided to redo the analysis every 6 month? I highly recommend the authors to read all the README-files for the various gene prediction datasets that GenBank and the other databases provide. E.g. Ensemble decided in 2012 (if I remember correctly) to not use any cDNA data anymore for validating their gene predictions. Thus, many exons with cDNA data available (e.g. from isolating and sequencing single genes by research groups) are not present in Ensemble’s gene predictions anymore, if these exons are not predicted by gene prediction tools. Similarly, there are many predicted exons for which no evidence (cDNA/RNA-seq) is available. Please check the GETx-project: There you can see how many RNA-seq reads in each tissue are found for each transcript. You will find out, that for many of the “alternative” transcripts, there is not a single read. The presented discussion of the variants (figures 3 to 6) supports my assumption that most predictions are just wrong. Could the authors provide any reference that it would be possible for such a seven-transmembrane-protein (the rhomboid proteases) to result in a functional protein if e.g. 1 transmembrane region in the middle would be missing due to alternative splicing? If 1 transmembrane region would be missing (this is suggested by several variants the authors discuss) this would turn the direction of the rest of the transmembrane helices and regions: what was inside before would be outside, and what was outside would be inside. To all what I know from membrane protein structures, the transmembrane helices stick together forming a dedicated tertiary structure. Could the authors provide a reference that transcripts with early terminations would ever result in stable and functional proteins, if e.g. only some of the 7 transmembrane helices are present anymore? The C.elegans ROM-4 was stated to contain a variant with just the first TM-helix present.
the authors really think that this would result in a functional protein? How do the authors exclude that such misspliced/unstable variants would not result in NMD?

=> The authors claim several times throughout the entire manuscript that they analysed selected model species without giving any information how these were selected. E.g. wikipedia contains a list of about 100 model organisms. Which was the rationale to just look at the 6 in the manuscript? Why not more plants? In this respect, there are many statements of the authors that are just wrong, such as “Currently, of the sequenced model genomes available,”. Of the 100 model organisms at wikipedia (and there are likely more model organisms if other researchers were asked) at least for 80 of them complete genomes are available. Altogether, there are about 5000 eukaryotes with genome assemblies available, of which at least 4000 are complete (e.g. check www.diark.org). In this respect, all the speculations and discussion about which organism contains the most homologs or the most variants are just speculations and should explicitly termed as such. E.g. Brassica species underwent another whole-genome-duplication after separation from the Arabidopsis, and will thus contain many more than the 22 Arabidopsis homologs. Fish also underwent another 1 or 2 (depending on lineage) whole-genome-duplication, therefore will also contain more homologs.

=> The authors state several times that, although not observed, they expect more alternative variants for the Arabidopsis homologs, because many more variants were identified for human. What is the basis for this expectation? Is there any reference that demonstrates, that Arabidopsis genes have as many alternative splice variants as mammalian genes, on average? All what I am aware of just contradicts this expectation. Arabidopsis genes have fewer exons (thus less possibilities for alternative splicing) and less alternative splice events. Why do the authors not expect more alternative variants for Drosophila or C.elegans, based on their rationale? Shouldn't yeast have at least some variants (doesn't have any yet)? Of course, yeast doesn’t, but this should make the authors aware of the problems in their argumentation.

=> Although the manuscript deals with alternative splice variants, I did not find any reference on the ample literature on this subject. Not even references to a few reviews. Similarly, there is not even mentioning of the accepted types of alternative splicing, e.g. differentially included exons, alternative 5'/3’ splice sites, mutually exclusive splicing, etc. Categorizing the variants detected with these categories would be much more informative than categorizing by region. Based on the descriptions in the supplementary tables, many variants are highly likely just the result from sequencing errors leading to frame-shifts or alternative amino acids. There is only a single alternative splice form that would lead to alternative amino acids for a certain regions, which is mutually exclusive splicing, but this information I did not find anywhere. This could easily be confirmed.

=> I did not see any gene structure in the manuscript or supplements. The usual procedure in the field is to provide a gene structure drawing of each gene and mark the splicing events on these structures. This is common practice since >30 years. The authors do not show any protein sequence, nor any cDNA sequence, nor any alignment. I cannot see any use of the provided tables in the supplements for other researchers. Terms such as “Original UTR missing, new UTR generated from coding region” (table S3) are not useful. How can a UTR be generated from a coding region? Does this mean that this is just an alternative translation start site? What does “Default 5’UTR missing, extended downstream from isoform 1” mean? When I think of a gene structure with exons and introns, which splice event would represent this prosaic description? I will not provide more examples here, but by browsing through the supplementary tables I did not find a single useful description. Thus, the authors should provide a gene structure scheme for each gene and mark the events for each gene accordingly. Such schemes would represent exact descriptions of the splicing events. All prosaic descriptions need to be removed.
The authors state in the Methods section: “This was necessary to acquire sequences that resulted from alternative splicing only, as opposed to derivations from other routes.” Which other routes would lead to variants, if not alternative splicing?

The authors state in the Methods section: “A prime example of such a situation occurred with the human database entries, where the newest version contained more verifications than the previous versions.” How did the authors determine, whether the newest version contains “more verifications”? Is there any reference that GenBank obtained more full-length cDNA data in 2016 and that these data were used in the gene predictions?

The authors confirm by experiments the alternative splicing of two Arabidopsis genes, but exactly these variants have already been published by the same group some years ago. Thus the authors do not provide any further and new evidence that any of the predicted variants of the other genes is present and functional. The authors would not even need to do experiments to show functionality. If they only analyse those transcripts for which cDNA/EST data are available, these cDNA/EST data are already the evidence. Further evidence could be obtained if the same transcripts were found for closely related species.

The Introduction is extremely unbalanced. E.g. for this very general statement “Rhomboids are also part of an even larger group of proteins involved in regulated intramembrane proteolysis 2–31.” the authors provide 30! references, but the authors miss to provide a useful description of the protein superfamily in general. As far as I understand, these proteins are present in bacteria, archaea and eukaryotes. It would be essential information how the various subgroups are phylogenetically separated. Without such information, there is no possibility to understand the variants. For example, which of the Arabidopsis homologs are the result of the many plant whole-genome duplications? Do such duplicates have identical alternative splicing? Similar for human and mouse: Which homologs are related to which Drosophila and C.elegans homologs, do the orthologs in human/mouse (derived by the two whole-genome duplications at the origin of the vertebrates) have identical splice variants?

Is the work clearly and accurately presented and does it cite the current literature?  
Partly

Is the study design appropriate and is the work technically sound?  
No

Are sufficient details of methods and analysis provided to allow replication by others?  
No

If applicable, is the statistical analysis and its interpretation appropriate?  
Not applicable

Are all the source data underlying the results available to ensure full reproducibility?  
Partly

Are the conclusions drawn adequately supported by the results?  
No

**Competing Interests:** No competing interests were disclosed.
I have read this submission. I believe that I have an appropriate level of expertise to state that I do not consider it to be of an acceptable scientific standard, for reasons outlined above.

Author Response 04 Apr 2018
Kenton Ko, Queen's University, Kingston, Canada

Thank you for taking the time and effort to review our submission. This is very much appreciated!

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We will be in touch soon with the next version.

**Competing Interests:** None

Author Response 22 May 2018
Kenton Ko, Queen's University, Kingston, Canada

Dear Dr. Kollmar,

Thank you for agreeing to review version 1 of our submission and helping us with your assessment. This is much appreciated!

We now have version 2 available for your further assessment. Based on the feedback, Version 2 was revised substantially to address the aspects raised and to enhance clarity for the readers in the areas of writing style and additional details.

As per your suggestions, improvements were made to the presentation by removing references that may not be necessary to the readers. Removal of these extra references allowed us to revise the Abstract, Introduction, and Methodology sections to enhance clarity with respect to the study’s focus or purpose. The title was also modified slightly to reflect these revisions.

We have revised the Methods section substantially to clarify the approaches used and to provide more needed details concerning the analysis of available database entries and the functionality assays using recombinant proteins. Revisions were also made to the Source Datasets and the Supplementary Tables to enhance their presentation, to provide more descriptive titles and related aspects. We hope that there is now sufficient methodological details for others to follow.

The style of writing and presentation in the Methods was revised substantially. We have removed unneeded information and added needed details/information on the tools used, tools that reflect what was being done in the comparative analysis of available entries – namely that we compiled all available entries and assessed their status as splice variants using RNA and protein data. We have added information in the text and in the tables to clarify what the compiled entries represent, e.g., entries or accession numbers listed could represent predicted entries but they are listed and used as reference points after assessment with RNA and protein data.

We have added more information (or information deemed missing) concerning the recombinant
protein work and revised the information to enhance clarity. For splicing information pertaining to
the At1g74130 (L, M, S) and At1g25290 (L, S) protein variants used later in the study, we have
added citations and direction to the Supplementary Material for protein information.

The multi-year facet of the database work was removed to avoid confusion and to enhance clarity.
This removal allowed us to streamline the database analysis protocols and provide more details of
the tools used. The Results and Discussion sections were modified similarly to reflect these
changes. The removal of the multi-year aspect in combination with the above revisions should
help the readers interpret the work more clearly.

Concerning the aspect of protein variants with substantial changes being potentially functional, we
have provided citations as possible indications that major changes could still give rise to rhomboid
proteins with functionality, such as early termination resulting in proteins with one missing
transmembrane region.

The choice of model organisms used was explained better by revising the style of
writing/presentation and by characterizing the work as a comparative analysis or survey of
available entries, as opposed to a direct comparison of splice variant numbers between species
and its interpretation in this way. We hope that we have interpreted this suggestion correctly. This
aspect, we believe, was also raised by Reviewer 3, Dr. Orlov.

We did not include references dealing with alternative splicing because the study was not intended
to be at this level, but working at the level of surveying and comparing available entries and what
these entries may reflect at the protein level. As described above, substantial revisions were
focused on clarifying the intent of the work to avoid this impression. This reasoning applies to why
gene structures and splicing events were not provided. All of the supplementary tables were
revised to enhance the writing style of the descriptions and render them more useful. We hope
that the revisions provided clarity on these issues.

The latter comments concerning passages in the Methods section were addressed by the
revisions outlined above. This would also be the case for the comments pertaining to the
Introduction – addressed by the revisions outlined above.

Sincerely,

Kenton Ko and Josh Powles

**Competing Interests:** No competing interests were disclosed.
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