Optimization of skeletal protein preparation for LC-MS/MS sequencing yields additional coral skeletal proteins in *Stylophora pistillata*

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Abstract

Stony corals generate their calcium carbonate exoskeleton in a highly controlled biomineralization process mediated by a variety of macromolecules including proteins. Fully identifying and classifying these proteins is crucial to understanding their role in exoskeleton formation, yet no optimal method to extract and isolate and characterize coral skeletal proteins has been established and their complete composition remains obscure. Here, we tested four skeletal protein extraction protocols using acetone precipitation and ultrafiltration dialysis filters to present a comprehensive scleractinian coral skeletal proteome. We identified a total of 60 proteins in the coral skeleton, 44 of which were not present in previously published stony coral skeletal proteomes. Extracted protein treatment protocols carried out in this study revealed that there is no “one optimal method” and each protocol revealed a unique set of method-exclusive proteins. To better understand the general mechanism of skeletal protein transportation, we further examined the proteins’ gene ontology, transmembrane domains, and signal peptides. We found that transmembrane domain proteins and signal peptide secretion pathways, by themselves, could not explain the transportation of proteins to the skeleton. We therefore propose that proteins are transported to the skeletal via vesicles and possibly non-traditional secretion pathways.
Keywords
Skeleton organic matrix proteins (SOMPs), biomineralization, Scleractinia

Introduction
Scleractinian corals, or stony corals, are the most prolific biomineralizers in phylum Cnidaria [1]. They are a key components of shallow-water tropical reefs, often forming massive structures that serve as the foundation of an ecosystem which hosts some of the more biodiverse communities on the planet [2, 3]. They are amongst the oldest biomineralizing metazoans, producing calcium carbonate (CaCO₃) exoskeletons in the form of aragonite through biologically-directed mechanisms [4, 5], which makes up $\geq 95\%$ of the entire skeletal mass, the remainder of which is the skeletal organic matrix (SOM) [6-8].

Biomineralization refers to the ability of a living organism to selectively exploit elements from its surrounding environment to build a biologically functioning crystalline structure [9, 10]. Biomineralizers can be found throughout all kingdoms of life, from bacteria [e.x. 11], to algae [e.x., 12], mollusks [e.x., 13], corals [e.x., 10] and mammals [e.x., 14]. The minerals formed through this process differ in structure from their non-biological counterparts and their formation is mediated by a variety of organic molecules (the SOM), which have been intensively studied since the 1960's in diverse organisms [reviewed by 10]. The SOM consists of proteins, lipids, and polysaccharides, which are not present in the geological mineral form [reviewed by 15]. SOM proteins (SOMPs) embedded within the mineral are hypothesized to serve as a framework for crystal nucleation [reviewed by 16]. The proteins involved in the process of skeletal biomineralization have been described and studied extensively in echinoderms [e.x., 17, 18] [and reviewed by 19, 20], mollusks [e.x., 21, 22-24] and
mammals [e.g., 25, 26, 27], among others [28]. At present, the best described SOMP complex is of mammalian bone and teeth [15]. In stony corals, the most current knowledge of SOMPs is limited and based on intraskeletal protein extraction [29-33]. It has been suggested that coral SOMPs aid in the molecular processes of crystallization as well as in the development and strengthening of the minerals ([34-37] among others). The constant advancement of mass spectrometry technology has broadened our capability to identify many proteins in skeletal extracts, even those proteins in low abundance [38]. However, this technology is sensitive to contamination by organic matter remnants from soft tissue and cell debris from the study organism, and the little-addressed issue remains of contamination by researchers during the protein extraction, preparation, and sequencing steps [39-41]. Upon extraction, the SOMPs are usually divided into two fractions: acid-soluble matrix and acid-insoluble matrix proteins (ASM and AIM respectively), based on their solubility in the acid of choice [22, 31-33, 42]. While some past attention has been directed towards the soluble fraction [30, 43, 44], Pereira-Mouriès et al. [45] showed that, in the bivalve *Pinctada maxima*, the classification of AIM and ASM is misleading and that both fractions share common features. Furthermore, Goffredo et al. [42] found in the stony coral *Balanophyllia europaea* that both fractions consist of the same macromolecules; they associated the degree of solubility to the difference in cross-linking. They also showed that each solubility fraction has a different influence on calcium carbonate crystal morphology, aggregation, and polymorphism in-vitro. In contrast, Ramos-Silva et al. [32] observed a different SOMP composition between solubility fractions in the scleractinian coral *Acropora millepora*. Out of 36 SOMPs, only two were found exclusively in the soluble fraction and twelve were exclusive to the insoluble fraction. These examples
demonstrate the attempts to attribute different properties to the two fractions but the data remain inconclusive.

To date, three major coral skeletal proteomes have been published [31-33] with each proteome consisting of 30-40 proteins. Of the 30 proteins sequenced from *A. digitifera* skeleton, 26 were also detected in *A. millepora* skeleton [32, 33]. They consist mostly of either transmembrane (TM) domain proteins or secretory proteins [33]. However, only 12 of the proteins identified in *A. millepora* skeleton matched those found in *S. pistillata* skeleton [31, 32]. In *A. millepora*, 11 TM domain-containing proteins were identified, as well as two proteases that were not detected in *S. pistillata* [32]. The authors suggested that the proteases’ role is in cleaving the extracellular domain of TM proteins and incorporating them into the skeleton.

The coral skeletal proteomes published to date reveal an overlap of several detected proteins, but at least 10 proteins from each species appear to be unique. It is currently unknown if this is truly due to species-specific gene expression and protein localization or to methods in extracting, purifying, and sequencing the proteins. In this study we analyzed several methods for protein purification to increase the detection of proteins from cleaned coral skeleton powder. We show that the use of acetone precipitation versus centrifugal filter washing, and the degree to which each purification method is performed, affects the numbers and types of proteins that can be sequenced by mass spectrometry. Further, we suggest that there is no one ‘best’ method for coral skeletal protein preparation such that future research projects may need to utilize several preparation methods to detect the full breadth of proteins embedded in coral skeleton.

**Methods**

*Sample collection and preparation for protein extraction*
The hermatypic coral *Stylophora pistillata* (Esper, 1797) was collected under a special permit from the Israeli Natural Parks Authority in the waters in front of the H. Steinitz Marine Biology Laboratory, Eilat, Israel, Red Sea (29°30 N, 34°56 E), using SCUBA diving.

We fragmented one *S. pistillata* colony into small pieces, approximately 2x2 cm, with a diamond band saw. Coral fragments were transferred to 50-ml Falcon brand conical vials (Falcon tubes) and oxidized with 20 mL 1:1 of 30% H$_2$O$_2$: 3% NaClO solution for 1 hour, during which 1.5 mL of 3% NaClO solution were gently added to the tubes every 20 minutes and continued the incubation overnight at room temperature following modified methods of Stoll et al. [46]. Fragments were washed with double distilled water five times for one minute each time and dried at 60°C overnight. We crushed the cleaned fragments to ≤63 µm diameter with a mortar and pestle. Skeleton powder was then oxidized in sterile Falcon tubes and washed as above three more times (i.e., four complete rounds of oxidative cleaning) to ensure that no organic residue remained on the skeletal grains. In each cycle, the removal of the oxidizing or washes solution was performed by centrifugation at 5,000 x g for 3 min at 4°C. Cleaned skeletal powder was then dried overnight at 60°C. We carried out all the described processes in a laminar flow biological hood (apart from oven drying) with all preparation tools and surfaces bleached to avoid contamination.

To monitor the removal of proteins from the skeletal fragments, we checked the cleaning efficiency under SEM after the fourth oxidative cleaning. Samples were vacuum-coated with 4 nm gold prior to examination under a ZEISS Sigma TM scanning electron microscope an in-lens detector (5kV, WD = 5-7mm) (SI Figure 1a,b). In addition, we sonicated the powder at 4°C in filter-sterilized phosphate buffered saline (PBS, pH 7.4) for 30 minutes, pelleted the powder at 5,000 x g for 3
minutes at 4˚C, concentrated the supernatant on a 3-kDa cutoff centrifugal filter unit (Amicon) and loaded samples of supernatant on a 8-16% SDS-PAGE TGX stain free gels (Bio-Rad) (SI Figure 1c).

Extraction of skeletal proteins

We decalcified approximately 1.3 g cleaned skeleton powder per protein extraction protocol in 0.5 M acetic acid (30 ml acid/g cleaned skeleton powder) in Falcon tubes while rotating the tubes at room temperature for 3 hours. Samples were then centrifuged at 5,000 x g for 5 min at 4˚C and supernatant was transferred to a new tube and stored at 4˚C. We continued the decalcification of the undissolved pellets with a second volume of 0.5 M acetic acid and allowed decalcification to proceed to completion. We then combined both liquid fractions (70 ml total) for each sample, froze the total volumes at -80˚C, and dried them by overnight lyophilization. The dried pellets were stored in -80°C until further processing.

The lyophilized pellets were re-suspended in 12 ml MilliQ water and the proteins were concentrated on 3 kD cutoff Amicon® Ultra 15 centrifugal filter units (Merk-Millipore) 5,000 x g at 4˚C to reach a final volume of 0.5 ml. To continue desalting the samples we diluted them again to 12 ml in MilliQ water and repeated the concentration procedure as above. At this stage, we observed an insoluble pellet in all samples.

Because washes of the extracted pellet can reduce the representation of proteins, in this study we examined two protein concentrating and cleaning methods performed for previously published coral skeletal proteomes; (i) centrifugation ultrafiltration (CF methods [32]) and (ii) acetone precipitation (ACT methods, [31]) (Table 1; SI Table 1). Further, for each method we separately examined the ASM and AIM fractions.
Two filter-concentrated samples containing both soluble and insoluble fractions were examined for the effects of centrifugal filtration. Sample CF2 was centrifuged at 5,000 x g for 5 min at 4°C to pellet the AIM fraction. Both the ASM and AIM of CF2 were divided into separate sterile 1.5 ml microcentrifuge tubes, lyophilized, and stored at -80°C until further use. Sample CF4 was desalted as described above for a third time before separating ASM and AIM fractions.

Both ACT samples were centrifuged as above for CF2 to separate ASM and AIM fractions. To sample ACT1 ASM (concentrated to 0.5 ml by centrifugal filtration as described above) was added 2 ml 100% ice cold acetone. The sample was vortexed for 10 seconds, incubated at -20°C for 30 minutes, and centrifuged at 4,300 x g for 30 min at 4°C. Finally, the pellet was washed three more times with 2 mL of 80% ice cold acetone and stored then at -80°C until further use. The AIM fraction was similarly washed four times with 80% acetone. Both fractions of sample ACT3 were treated as in ACT1 but with one less washing step of each fraction.

**LC MS/MS**

*S. pistillata* skeletal protein samples were dissolved in 5% SDS and digested with trypsin using the S-trap method overnight at room temperature. We analyzed the resulting peptides with nanoflow ultra-performance liquid chromatograph (nanoAcquity) coupled to a high resolution, high mass accuracy mass spectrometer (Fusion Lumos). The sample was trapped on a Symmetry C18 0.18*20mm trap column (Waters, Inc) and separated on a HSS T3 0.075*250 mm column (Waters, Inc.) using a gradient of 4-28% (80% acetonitrile, 0.1% Formic acid) for 150 minutes. Spray voltage was set to +2kV. The data were acquired in the Fusion Lumos using a Top Speed Data-Dependent Acquisition method using a cycle time of 3 s. An MS1 scan was performed in the Orbitrap at 120,000 resolution with a maximum injection
time of 60 ms. The data were scanned between 300-1800 m/z. MS2 was selected using a monoisotopic precursor selection set to peptides, peptide charge states set to +2-+8 and dynamic exclusion set to 30 s. MS2 was performed using HCD fragmentation scanned in the Orbitrap, with the first mass set to 130 m/z at a resolution of 15,000. Maximum injection time was set to 60 ms with automatic gain control of 5x10^{-4} ions as a fill target. The resulting data were searched against the NCBI *Stylophora pistillata* protein database using the Byonic search engine (Protein Metrics Inc.) – the first search was carried out without any false discovery rate (FDR) filtering, to generate a focused database for a second search. The second search was set to 1% FDR, allowing fixed carbamidomethylation on C and variable oxidation on MW, deamidation on NQ and protein N-terminal acetylation. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the Pride [47] partner repository under the dataset identifier PXD017891; reviewer login username and password have been provided to the reviewers. Data will be made publicly available without a required login in ProteomeXchange upon publication of the manuscript.

**Data sorting**

We used the *S. pistillata* genome database as a reference peptide database for the mass spectrometry analysis [48] (NCBI BioProjects PRJNA281535 and PRJNA415215) appended with known NCBI *S. pistillata* skeletal proteins[e.x., 31, 49] *S. pistillata* carbonic anhydrase, ACE95141.1). We also included a common contaminants database. Despite this inclusion of contaminants databases, several proteins likely of human origin were sequenced and attributed to *S. pistillata*. To filter out these potential contaminants from our final list of coral-specific proteins, we BLASTed all sequences against the ‘Primates’ database in NCBI using Blast2GO.
We then examined NCBI-generated sequence alignments of coral versus *Homo sapiens* proteins with e-values lower than e\(^{-50}\) and percent mean similarity greater than 50%, all sequences with e-values lower than e\(^{-100}\), and all sequences with percent similarity greater than 80%, and removed from our final list of coral proteins any sequences with three or more peptides each of seven or more amino acids in length that were identical between *S. pistillata* and humans.

All proteins identified by the LC MS\text/MS analysis were filtered to proteins with at least two significant spectra and at least one significant unique peptide. Skeletal proteins were first sorted by fractions and methods (ASM and AIM for each extraction method). Next, we sorted all skeletal proteins by their gene ontology (GO) terms [50, 51] to differentiate plasma membrane proteins and extracellular matrix (ECM) proteins. Finally, we grouped all by terms of interest: protein modification, transmembrane, and ECM; membrane processing; metal binding and vesicular.

Proteins detected by LC-MS/MS were annotated using the Trinotate pipeline which relies on both pfam and Uniprot data [52]. GO terms and PFAM annotations were assigned to *Stylophora pistillata* predicted proteins and transcripts using Trinotate 3.0.1 (https://github.com/Trinotate). Transmembrane regions were predicted using the TMHMM server v2.0 [53]. Signal peptides on the N-termini of proteins were predicted using SignalP 5.0 [54, 55]. Attachment of proteins to the exterior of the cell membrane by glycosylphosphatidylinositol (GPI) anchors was predicted using PredGPI [56]. These computational analyses used program default settings and cutoffs. Completeness of protein sequences was determined by comparing all returned coral proteins to the *Acropora digitifera* genome [57]; NCBI RefSeq assembly GCF_000222465.1). Detection of various proteins across solubility and protocol fractions was visualized in Venny 2.1 online software.
We determined orthologous biomineralization genes, known from skeletal proteomic analysis, across coral taxa. First, we estimated orthology relationships between all non-redundant genes of selected metazoan species using OrthoFinder. We included all genes of all Cnidaria species with known genome-based annotations. OrthoFinder generates orthology-groups (Orthogroups) based on normalized reciprocal best Blast hits’ bit scores [58], and then estimates orthologues genes pairs within Orthoroups [59]. We then selected all pairs of *Acropora* spp. orthologs to *S. pistillata* (1:1, 1:many, many:many relationships) ([31-33], this study). From these pairs we further selected *S. pistillata* spectra-based identified proteins, or skeletal proteins known in the literature. Since not all skeletal protein annotations from the literature were included in our reference Orthofinder proteome datasets, we further found their best matches in the reference orthofinder proteome using blastp.

**Results:**

After extensive cleaning of the powdered skeleton and acid-extraction of embedded organic matter we identified in total 60 coral-specific proteins, with two or greater spectra or one unique peptide with at least 10 spectra and an identification score of 250 or greater, in *S. pistillata* skeleton as predicted by the species genome [48] (Table 2, SI Table 2). Trinotate [60] [52] returned GO annotations [50] for all of these proteins, although many remain uncharacterized (SI Table 3).

In order to evaluate the efficacy and improve current methods for stony coral skeletal protein extraction, we examined four different protocols; two centrifugation ultrafiltration filters (CF) and two further acetone precipitation (ACT) protocols. Proteomes of the four methodologies differed in composition and variety (Figure 1). Combining results of all CF fractions identified 52 coral-specific proteins while combined ACT protocols yielded 13 such proteins (Figure 1A). Moreover,
redundancy between methodologies was low. Only 8.3% of the proteins overlapped between methods while 78.3% and 13.3% of the proteins were exclusive to combined CF and combined ACT fraction data, respectively.

To evaluate the extracted protein treatment efficiency of each protocol, we first compared the number of proteins detected in each method (e., ACT1 vs ACT 3 and CF2 vs CF4). Of the 8 proteins found only in samples from the acetone wash protocols (ACT 1 and ACT 3), one was observed only in ACT1 while five were observed only ACT3 (Figure 1, SI Table 2); both ACT1 solubility fractions had one more wash step than did those in ACT3. In contrast, of the 47 proteins observed only in CF samples, 11 were found in CF2, which went through two filter centrifugation steps, while 16 were found in CF4, which went through a third filter centrifugation step. Further, only two proteins were observed in all four purification fractions in at least one solubility form: CARP4/SAARP1 and synapsin 2-like. Two additional proteins were observed in three of the four protocols: STPCA2 and an uncharacterized protein.

We next examined the difference in protein composition obtained from the solubility fractions ASM versus AIM. We identified different distributions of proteins in the ASM vs. AIM in both purification methods (Figures 1B & C). Notably, when combining all methods, more proteins were identified in the AIM compared to the ASM. A total of 57 coral proteins were identified in the AIM fraction compared to a total of 8 in the ASM fraction (SI Table 2), of which 5 (8.3%) were identified both in ASM and AIM.

Our data compared to other coral skeletal proteomes

Since our analysis yielded a large amount of new skeletal proteins, we also compared our results with the three previously published proteome of S. pistillata, A. digitifera...
and *A. millepora*. Out of our entire identified skeletal proteome containing 60 proteins, using OrthoFinder and BlastP, only 16 were found to be similar to proteins identified in these studies. Yet, this proportion of overlap (16 out of 60) is significantly greater than the expected proportion by chance, since the proportion of known skeletal matrix proteins in the reference coral proteomes is extremely small (less than \(~0.2\%)). Seven proteins were found to overlap all four proteomes: a coadhesin-like protein, an EGF and laminin G domain-containing protein, a hypothetical protein, a MAM and LDL-receptor class A domain-containing protein, a mucin, aspartic acid-rich protein 2-like, and a ZP domain-containing protein (Table 3).

**Skeletal proteome characterization**

We interrogated the mechanisms by which proteins may be exported from or attached to the cell (SI Table 2). Seventeen coral skeletal proteins with likely complete N-terminus predictions possess signal peptides as a potential mechanism for export from the cell. Eight proteins contain at least one transmembrane span suggesting that they are embedded in the cell membrane. Further, 10 proteins likely interact with the exterior of the cell membrane by GPI anchors. In total, 25 of the 60 sequenced coral skeletal proteins exhibit documented characteristics for localization in the ECM.

Because the majority of the proteins sequenced from the *S. pistillata* skeleton do not possess features for signaling their export from the cell, we queried the data set for further suggestions of positioning the proteins in the membrane or that the proteins may be exported by vesicles such as those that may be involved in calcium concentration. To do this, we examined the skeletal proteome annotations and GO classifications toward finding common features to allow grouping of proteins. Out of the entire skeletal proteome sequenced in this study (60 proteins), 39 genes were
returned with GO terms that allowed their classification into five groups of interest based on their cellular component, biological process, and molecular function to suggest likely cellular locations pertinent to the calcification mechanism, which may therefore be indicative of their function in this process: lipid\phosphate\glycan related proteins (i.e., membrane processing); ECM-related, transmembrane, and protein modification proteins; and metal binding proteins vesicular/secretion related proteins, (SI Table 3).

Of the proteins with GO terms, 10 proteins are suggested to be involved with processing of the cell membrane (Figure 2), A much larger number are related to vesicles/secretion as well as binding metal, with 21 and 25 assigned to each category, respectively. We combined these two categories in our proposed cellular location Figure 2 as some of the skeletal proteins proposed to be found intracellularly in vesicles are also known to bind calcium [34, 61]. Finally, 19 and 12 are potential ECM proteins or are involved in protein modification, respectively. It should be noted that many protein are assigned to multiple categories.

Discussion:

In this study we show the importance of using complementary post-extraction methods to concentrate coral skeletal proteins for sequencing the full breadth of the skeletal proteome. Our results show a clear and marked difference in detected proteins between protein purification methods, with only two proteins observed in all methods. Centrifugal Filter (CF) methods yielded a much greater abundance and diversity of proteins than did acetone precipitation (ACT) (Figure 1). Moreover, we found that protein detection is largely method-exclusive (Figure 1). We hypothesize that the protein yield differences between extraction methods is based on the different properties of each method. CF is a mechanical filtration based on size and has a bias
toward hydrophobic proteins, whereas ACT is based mainly on the chemical
interactions of proteins resulting in increased precipitation of hydrophilic proteins
[62]. We speculate that the CF methods yielded more total proteins in our study as
acetone precipitated proteins may be difficult to resolubilize, potentially leading to the
possible loss of many proteins in the pellet which is not transferred to the trypsin
digestion step.

A major challenge in working with intra-skeletal proteins is isolating the true
skeletal proteins from soft tissue contamination [40, 41]. In the present study, as in
previous work on S. pistillata skeletal proteins [40] we carried out an intensive
oxidative cleaning step on the skeletal powder, in addition to cleaning the skeletal
fragments, to avoid contamination. We did not observe any organic residues by SEM
on intact skeletons following the second cleaning step (SI Figure 1a,b) or in PBS
soaked on cleaned powders and concentrated (SI Figure 1c), and we are therefore
confident that all sequenced proteins are endogenous to the skeleton and are not coral
cellular contamination.

In this study, we examined the protein composition of the ASM and AIM
proteins separately. Although we see variations due to the different purification
techniques (Figure 1), our results indicate that, broadly, the AIM is distinguished in
composition from the ASM. Across all purification techniques, the AIM fractions
yielded more total proteins and fraction-exclusive proteins than the ASM fraction did
(Figure 1B, C). Since all extraction and precipitation experiments were carried out
simultaneously, we can rule out batch effect and technical differences. This AIM
versus ASM skeletome fractional composition is similar to previous studies carried
out on other marine biomineralizers such as mollusks [45] as well as to the stony
corals A. millepora [32] and S. pistillata [31]. However, the overlap between the two
solubility fractions reported for *A. millepora* [32] was 61%, in contrast to our study which shows an overlap across all purification treatments combined of only 10% (Figure 1). One plausible explanation for this difference is the use of improved LC-MS/MS performance over the past several years enabled us to detect more proteins and obtain a more comprehensive proteome. In previous studies of coral skeletal proteomes which detailed differences between solubility fractions, the instruments used were low resolution, low mass accuracy, which tend to result in a lower percent identification of the data. Additionally, most search engines and FDR-calculating algorithms struggle with very small datasets [63, 64]. Our use of Byonic helps to alleviate statistical limitations that can result in false negative results.

Further differences in proteomes, beyond species differences, are the differing reagents used in precipitation (compared to [31]) and our smaller centrifugal filter cutoffs (compared to [32]). This second difference is particularly important with respect to protein degradation. Even when embedded in biominerals so that amino acids and even short peptides persist, proteins may still succumb to degradation [65]. This result in small peptides fragments that may be lost from centrifugal filter units with 10 kDa pore sizes and larger. It is reasonable to assume that coral skeletal proteins go through the same process, and if so, the cutoff of the membrane directly affects the number of peptide spectra matches (PSM). Using smaller cutoff filters in this study might have allowed us to capture some of these sheered peptides and led to higher PSMs.

Traditional protein trafficking by signal peptides and transmembrane domain does not explain the full extent of protein transportation to skeleton. Corals’ skeleton is external to the animal; therefore, proteins in the skeletal matrices must be transported outside the cells or span the membrane and have an extracellular
portion to reach the skeletal crystallization front. Indeed, recent studies of anthozoans reveal a significant proportion of TM domain proteins (~35%) in the SOM [32, 33], similar to that found in the better-studied Echinoidia [66]. Based on these findings, we examined the hypothesis that many of the skeletal proteins originate in the plasma membrane.

Our analysis revealed that TM domain proteins are not the major component of the SOM protein complex; in the present proteome, TMHMM prediction suggests that eight are embedded in the membrane. Hence we examined a cellular secretion option. Ramos-Silva et al. [32] reported 15 proteins (41%) with SPs that did not also possess a TM domain in the A. millepora proteome. In their study of the A. digitifera proteome, Takeuchi et al. [33] reported a similar proportion of proteins with SPs but no TM domains (40%). When examining the combined S. pistillata proteomes, 17 out of 60 (28%) were positive for SPs (SI Table S2, [31]), likely a combination of incomplete gene prediction and other mechanisms for exporting the proteins to the calicoblastic space.

Hydrophobic regions of TM domains are similar to those of SPs, making SP prediction difficult [53, 67]. We therefore cannot say for sure, based on these two analyses (TMHMM and SignalP) alone, which is the more dominant pathway for protein transport. While SP by itself can help predict traditional secretory pathways [67], proteins that do not possess SPs may still leave the cell via other means without classic SPs. This includes membrane pores [67], ATP-binding cassette transporters [68] and autophagosome/endosome-based secretion [69, 70]. These are but a few examples of secretory pathways which bypass the conventional TM/SP signaling pathway (reviewed in [67, 70, 71]).
A further method by which protein may be exported to the site of calcification is a vesicular pathway that differs from the conventional SP and TM pathway. Previous studies have shown Ca\(^{2+}\) rich granules in the calicoblastic epithelium (skeletogenic cells), but not in the other tissue layers, suggesting their role as a Ca\(^{2+}\) reservoirs in the cells. Vesicles were previously identified in corals [72, 73]; however, their origin and content was not detailed and they are sometimes attributed to preservation byproducts. These intracellular ion-rich vesicles may endocytose sea water [74], after which they are enriched in carbonate ions and then form hydrated ACC and anhydrous ACC precursors stabilized by acidic biomolecules including CARPs [75]. Using cell cultures, Mass et al. [61] suggest that the vesicles, which contain Asp rich proteins, then transport their contents to the ECM, releasing their content by exocytosis. The biomineral then further develops extracellularly, likely aided by other ECM proteins [30, 35, 49, 76] as well as other biomolecules [77-80]. At present the process of calcium delivery to the skeleton and the role of most of the proteins in skeletal depositing remain to be determined.

Since TM and SP analysis did not fully explain the mechanism of protein transportation, we classified the likely functions and locations of the skeletal proteins according to their GO terms as lipid\(\backslash\)phosphate\(\backslash\)glycan related proteins, metal binding proteins, vesicular\(\backslash\)secretion related proteins, ECM-related and transmembrane proteins, and protein modification proteins (Figure 2, SI Table 3). Looking at the proposed locations or functions of skeletal proteins (Figure 2), we find comparable numbers of vesicular and metal binding proteins versus TM domain/ECM proteins. These results are in-line with the trend reported in previous coral skeletal proteomes [31-33]. While we do not exclude the role of TM proteins in the skeletal deposition
process, we suggest that the biological mechanism of SOM transportation is enhanced by vesicular secretion and other possible, non-traditional, secretion pathways.

It is difficult to map and characterize proteins in non-model organisms such as *S. pistillata* [81-83]. High quality proteomic mapping requires knowledge of phosphorylation, glycosylation, proteolytic cite activities and other modifications [31, 84-86], in order to create a more thorough database [81-83]. Further, many of the coral skeletal proteins reported to date remain uncharacterized. Uncharacterized proteins were reported in *Acropora* skeletons at a rate of approximately 25% [32, 33] while, they were previously reported at less than 10% in *Stylophora* skeleton [31]. Our study revealed a much greater proportion of 28% uncharacterized proteins. While partially attributed to sample size, it is most likely due to quality of genomic data available, since stony corals are non-model organisms and their genomic libraries are far from being complete, resulting in incomplete databases on which to map the proteome and many uncharacterized genes.

**Conclusion:**

In this study we have considered the differential effects of coral skeletal protein extract preparation as well as the method by which these proteins, or parts thereof, are transported from intracellular to extracellular locations. When preparing coral skeletal proteomes, we propose that a multi-method approach to cleaning, demineralization, and protein extraction should be used. Our results showed that each protein preparation protocol yielded exclusive sets of proteins with little overlap between ACT and CF fractions. While CF protocols yielded many more proteins than did ACT methods, use of a single protocol to clean and concentrate coral skeletal proteins results in a significant amount of data loss, and it is therefore of crucial importance to consider alternative and complementary methods to obtain a fully comprehensive
skeletal proteome. We showed that while the role of TM domain proteins cannot be
overlooked, many of the proteins detected in the *S. pistillata* skeletal proteomes as
well as in that of other species point toward other secretory or vesicular pathways.
Our categorization method, supported by data from other recent studies, also suggests
that corals use an alternative secretory pathway, such as vesicles, and much work is
required in order to determine the calcium deposition pathway and the proteins
involved. Our study provides a large set of new uncharacterized coral skeletal proteins
as well as others of purported function but that have not been observed before in the
coral SOM. These data expand the current knowledge of the SOM in corals and will
help, in future studies, to resolve corals’ calcium deposition mechanism and the
various roles of the proteins involved.

**Abbreviations**

acetone precipitation of proteins (ACT)
acid soluble matrix (ASM)
acid insoluble matrix (AIM)
centrifugation ultrafiltration of proteins (CF)
false discovery rate (FDR)
gene ontology (GO)
glycosylphosphatidylinositol (GPI)
peptide spectra matches (PSM)
phosphate buffered saline (PBS)
skeletal organic matrix (SOM)
skeletal organic matrix proteins (SOMP)
transmembrane (TM)

**Declarations**
Ethics approval and consent to participate

Coral nubbins were collected under a special permit from the Israeli Natural Parks Authority.

Consent for publication

Not applicable.

Availability of data and materials

The datasets generated during the current study are available in the ProteomeXchange repository under file number PXD017891 (https://www.ebi.ac.uk/pride/archive). The reviewer login username and password have been provided to the reviewers. Data will be made publicly available without a required login in ProteomeXchange upon publication of the manuscript.

Competing interests

The authors declare that they have no competing interests.

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Author contributions

YP, JD, and TM designed the study. YP and RA prepared the samples. YP, RA, AM, ML, DM, and JD analyzed the data. All authors wrote the manuscript and approve this submission.

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**Table and Figure Legends**

**Tables**

Table 1: Summary of methods for protein concentration and cleaning after extraction in acetic acid. CF – Centrifugation filtration methods; ACT – acetone precipitation methods; ASM – Acid soluble matrix; AIM – Acid insoluble matrix.

Table 2: 60 coral skeletal proteins detected by LC-MS/MS across all treatments and solubility fractions. Proteins are listed in order of accession number. Gene ontology categorization is represented as aECM/transmembrane and protein modification, bmembrane processing, and cvesicle/secretion and metal binding.

Table 3. Orthologous coral skeletal proteins in the present work and previously published. Gene ontology categorization is represented as aECM/transmembrane and
protein modification, membrane processing, and vesicle/secrection and metal binding.

Figures

Figure 1: Distribution of proteins numbers according of the extraction methods.
Distribution of all proteins (SOM and ISOM combined) by extraction methods (A).
SOM and ISOM distribution by acetone precipitation methods (ACT). SOM and
ISOM distribution by centrifugation and ultrafiltration method (CF). Note that CF
methods yielded more proteins in total. ISOM Insoluble Skeletal Organic Matrix;
SOM Skeletal Organic Matrix.

Figure 2. Suggested cellular and extracellular locations of *S. pistillata* skeletal protein
based on GO terms. Large yellow and red circles represent calicoblastic cells with
red cell membrane facing the skeleton. Small green circles represent vesicles. And
purple squiggles represent transmembrane proteins that may or may not span the
width of the calicoblastic space plus ECM proteins.
| Sample Name                  | Filter desalting (CF) | Acetone Precipitation (ACT) |
|-----------------------------|-----------------------|-----------------------------|
|                             | CF2  | CF4  | ACT1 | ACT3 |
| Solubility Fraction         | ASM  | AIM  | ASM  | AIM  | ASM  | AIM  | ASM  | AIM  |
| Filter desalting x 2        | +    | +    | +    | +    | +    | +    | +    | +    |
| Filter desalting x 3        | -    | -    | +    | +    | -    | -    | -    | -    |
| Acetone precipitation       | -    | -    | -    | -    | +    | -    | +    | -    |
| 1x 80% acetone wash         | -    | -    | -    | -    | +    | +    | +    | +    |
| 2x 80% acetone wash         | -    | -    | -    | -    | +    | +    | +    | +    |
| 3x 80% acetone wash         | -    | -    | -    | -    | +    | +    | -    | +    |
| 4x 80% acetone wash         | -    | -    | -    | -    | -    | +    | -    | -    |
| gi Number   | Accession Number | Annotation                                                                 | Best score | Go Categorization | ASM | AIM |
|------------|------------------|---------------------------------------------------------------------------|------------|-------------------|-----|-----|
| 1263115389 | PXF12726.1        | Retrovirus-related Pol polyprotein from transposon 17A [Stylophora pistillata] | 340.80     | -                 | x   |     |
| 1263115517 | PXF12813.1        | hypothetical protein AW38_SpisGene23165 [Stylophora pistillata]            | 337.40     | -                 |     | x   |
| 1263116757 | PXF13377.8        | Sacin [Stylophora pistillata]                                             | 267.80     | -                 | x   |     |
| 1263117261 | PXF14205.1        | Proto-ecogene tyrosine-protein kinase receptor Rut, partial [Stylophora pistillata] | 336.70 | a,c               |     | x   |
| 1263119090 | PXF15740.1        | Protein FAM208A [Stylophora pistillata]                                    | 372.20     | -                 | x   |     |
| 1263119725 | PXF16398.1        | hypothetical protein AW38_SpisGene19330 [Stylophora pistillata]            | 273.10     | -                 |     | x   |
| 1263122270 | PXF18785.1        | Mucin-4 [Stylophora pistillata]                                            | 449.50     | a,c               |     | x   |
| 1263130525 | PXF26597.1        | Complement C3 [Stylophora pistillata]                                      | 283.10     | a,b,c             |     |     |
| 1263130510 | PXF26751.1        | Transmembrane protein serine 9 [Stylophora pistillata]                    | 244.20     | a                 |     |     |
| 1263131615 | PXF27832.1        | Poly [ADP-ribose] polymerase 11 [Stylophora pistillata]                   | 320.10     | c                 |     | x   |
| 1263136664 | PXF30831.1        | hypothetical protein AW38_SpisGene4366 [Stylophora pistillata]            | 322.20     | -                 |     | x   |
| 1263134737 | PXF39903.1        | hypothetical protein AW38_SpisGene4292 [Stylophora pistillata]            | 548.90     | -                 |     | x   |
| 1263135656 | PXF33181.0        | Nidogen-2 [Stylophora pistillata]                                          | 286.20     | a,c               |     | x   |

**gi Number**

**Accession Number**

**Annotation**

**Best score**

**Go Categorization**

**ASM**

**AIM**
| NCBI Accession Number | Annotation                                      | gi (NCBI) | Gene model id | RNA id | Acropora digitifera skeletal proteins (Takeuchi et al. 2016) | Acropora millepora skeletal proteins (Ramos-Silva et al. 2013) | Stylophora pistillata skeletal proteins (Drake et al. 2015) |
|-----------------------|------------------------------------------------|-----------|---------------|--------|-------------------------------------------------------------|---------------------------------------------------------------|-------------------------------------------------------------|
| ACE95141.1.a          | carbonic anhydrase                              | 190710633 | NA            | NA     |                                                             |                                                               | Sarasaparilla_g1749; mapped to ACE95141.1 (EU3532164) |
| PFX18785.1.c          | Mucin-4                                         | 126312270 | NA            | NA     | aug_v2a.098068.11                                           | JR987773                                                      | AGG36340.1                                                   |
| PFX30000.1            | hypothetical protein AWC38_SpatGene4292         | 126313473 | NA            | NA     | aug_v2a.063271.11                                           | JR972076.1                                                    | AGG36350.1                                                   |
| XP_02279720.1.b       | vitillogenin-like                               | 1270036141| gene3329      | mae552 | aug_v2a.088561.11                                           | aug_v2a.110868.11, JT001945.1                                 | AGG36347.1, AGG36349.1                                       |
| XP_02270690.1.a       | skeletal aspartic acid-rich protein 2-like      | 1270037961| gene4012      | mae498 | aug_v2a.014411.11, aug_v2a.014408.11, JT004496.1, JR991407.1| JR991407.1, JT004498.1                                        | AGG36358.1, AGG36357.1                                       |
| XP_02270694.1.c       | CUB and peptidase domain-containing protein 2-like | 1270037769| gene3981      | mae5452|                                                             |                                                               | JR970992.1                                                   |
| XP_02278239.1         | skeletal aspartic acid-rich protein 1-like      | 1270041141| gene5242      | mae762 |                                                             |                                                               | AGG36357.1                                                   |
| XP_022783415.1.a      | cadherin-like isoform X3                       | 1270043038| gene6099      | nai510 | aug_v2a.015945.11                                           | JT016638.1                                                    | AGG36341.1, AGG36335.1, AGG36343.1                           |
| XP_022780818.1.b      | major yolk protein-like isoform X2             | 1270045910| gene6661      | nai1721|                                                             |                                                               | AGG36343.1                                                   |
| XP_022789227.1.b      | haptoglobin-like protein                       | 1270052030| gene9579      | mae2971| aug_v2a.240151.11                                           | JT019463.1                                                    | AGG36343.1                                                   |
| XP_022794122.1        | galactoside-like isoform X2                    | 1270063049| gene13931     | nai823 | aug_v2a.186311.11                                           | JR976690.1                                                    | AGG36555.1, AGG36566.1, AGG36345.1                           |
| XP_022794736.1        | MAM and LDL-receptor class A domain-containing protein 2-like | 1270064196| gene14425     | mae9450| aug_v2a.099668.11, aug_v2a.099689.1, JT011118.1, JR994474.1| JT014391.1                                                    | AGG24391.1                                                   |
| XP_022790981.1        | uncharacterized skeletal organic matrix protein 8-like | 1270068394| gene16153     | mae2169 |                                                             |                                                               | AGG24391.1                                                   |
| XP_022790982.1        | uncharacterized protein LOC111333364           | 1270068396| gene16156     | mae2170 |                                                             |                                                               | AGG24391.1                                                   |
| XP_022804012.1.a      | EGF and laminin G domain-containing protein-like| 1270032917| gene23104     | mae2833| aug_v2a.063271.11                                           |                                                               | AGG36344.1, AGG36352.1                                       |
| XP_022806326.1.a      | ZP domain-containing protein-like              | 1270085816| gene23405     | mae30870| aug_v2a.076273.11                                           |                                                               | AGG36346.1                                                   |
| **ECM/Transmembrane & Protein Modification** | **Membrane Processing** | **Vesicle/Secretion & Metal Binding** |
|---------------------------------------------|-------------------------|--------------------------------------|
| Carbonic anhydrase 2-like                   | Complement C3           | Carbonic anhydrase 2-like             |
| Cation channel sperm-associated protein subunit beta-like | Endothelin-converting enzyme 1-like | Cation channel sperm-associated protein subunit beta-like |
| Chymotrypsin-like elastase family member 1 | Protein lingerer-like | Chymotrypsin-like elastase family member 1 |
| Coadhesin-like                             | Uncharacterized (6)     | Collagenase-like                      |
| Collagenase 3-like                         |                         | Complement C3                         |
| Complement C3                              |                         | Deleted in malignant brain tumors 1 protein-like |
| CUB and peptidase domain-containing protein 2-like |                         | Condensin-2 complex subunit D3-like |
| Digestive cysteine proteinase 1-like       |                         | Deleterious malignant brain tumors 1 protein-like |
| Endothelin-converting enzyme 1-like        |                         | Collagenase-like                      |
| EGF and laminin G domain-containing protein-like |                         | Endothelin-converting enzyme 1-like   |
| Hephaestin-like protein                    |                         | Low-density lipoprotein receptor-related protein 8-like |
| MAGUK p55 subfamily member 7-like          |                         | Mammalian ependymin-related protein 1-like |
| Major yolk protein-like                    |                         | Microtubule-associated tumor suppressor 1 homolog |
| Mammalian ependymin-related protein        |                         | Mucin-4                               |
| Mucin-4                                    |                         | Nidogen-2                             |
| Microtubule-associated tumor suppressor 1 homolog |                         |                                      |
| Nidogen-2                                  |                         |                                      |
| Proto-oncogene tyrosine-protein kinase receptor Ret |                         |                                      |
| Ras-like protein 3                         |                         |                                      |
| Skeletal aspartic acid-rich protein 2-like  |                         |                                      |
| Transmembrane protease serine 9            |                         |                                      |
| Uncharacterized (3)                        |                         |                                      |
| ZP domain-containing protein-like          |                         |                                      |

**Callicoblastic cell layer**

**Calcifying medium**

**Skeleton**