Research Article

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Miniaturized peptidomimetics and nanovesiculation in endothelin types through probable nano-disk formation and structure property relationships of endothelins’ fragments

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Abstract: Endothelins (ETs), which are multi-functional peptides with potential for antagonist-based-therapy in various physiological-malfunctionings, including cardio-vascular, nephrological, oncologic, and diabetic conditions, may produce newer chemical entities and drug leads. The present study deals with molecular-modeling of the ETs’ sub-types, ET-I, II, and III to find the structure property-relationship (SPR) of the ETs, and individual fragments derived from the ET sub-type ET-I. The ETs peptidic tails’ amino acid (AA) sequence’s structural differences and similarities, various dissected fragments of the ET-I, and SPR comparison with the sarafotoxin-6b (SRT-6b), a structurally-related snake-venom, showed points of dissimilarities for their structural specifications, geometric disposition, and physico-chemical properties. The generation of miniaturized (shortened sequence) peptides towards offering peptidomimetic compounds of near- and far-values compared SPR with estimations for log P, hydration energy, and other molecular and quantitative structure activity relationship (QSAR) were based on random and ordered-fragments derived from the original ET-I AA’s sequence, and sequential distance changes in the original ET-I sequence’s chain of 1–21 AA. The feasibility of alternate and bond length parameters-based possible cysteine–cysteine cyclizations, sequence homology, AA’s positional demarcation, and presence/absence of cysteines, homology-based basic non-cysteine and cysteines-AA based cyclization, total structure and fragments end-to-end cyclizations, and geometrical analogy-based miniaturized sequence of the shorter AAs from the original ET-I sequence, together with mutated replacements with naturally constituent AAs of the ETs, and SRT-6 sequences were utilized. The major findings of the fragmented sequences, and sequences at par with the

Graphical abstract: The present study deals with the molecular modeling studies of the endothelins (ETs), a multi-functional 21 amino acid (AA) peptides, with sub-types ET-I, II, and III to find the structure property relationship (SPR) of different fragments naturally derived from the cyclic body part of the ETs’ structures together with the structurally similar and common tail sequence (AA 16–21) utilization of the ETs and the dissimilar sarafotoxin-6b (SRT-6b) hexapeptide chain tail, as well as their SPR comparisons to find peptide-based leads as comparable to the ETs, especially ET-I. Approach to plausible vesiculation of the ETs and the involved process as a suggested mechanism is also discussed.

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original ETs to provide structures similar to the size, volume and with molecular and electronic properties of electrostatic potential and total charge density distribution, crucial factors in receptor bindings were investigated. The SPRs, molecular properties, and QSAR values were estimated to compare and validate the findings with the known homologous compounds, ET-I, and its known and potent antagonists. The study resulted in leads of smaller and larger sizes of peptide-based compounds which may have prospects as potent antagonist and in future needs their bioactivity evaluations after the synthesis. Moreover, approach to plausible vesiculation of the ETs, and the involved processes and structural requirements, together with the molecular interactions in settling a nano-vesicle of the peptidic structure with a possible mechanism is also suggested.

**Keywords:** endothelins, sarafotoxin-6b, structure property relationships, compartmentalization, miniaturized peptide, peptidomimetics, vesiculation

### 1 Introduction

The endothelin family of peptides, collectively termed as endothelins (ETs), comprises of endothelin-I (ET-I), endothelin-II (ET-II), and endothelin-III (ET-III), with binding to at least four different known endothelin receptors, i.e., ET₄₄, ET₁₁, ET₂₂, and ET₃₃ in the biological system [1,2]. The ETs have varying regions of distribution in biological sites [3], and are produced by vascular endothelium through pre-pro-ETs resulting in 39 amino acids (AAs) precursor, big ETs, which is transformed through activities of endothelin converting enzyme located on the endothelial cell membranes into various ETs (Figure 1) [4–7]. The ETs and its receptors have been cloned [8–10] and molecularly characterized [11]. Various peptidic and non-peptide receptor antagonists have also been designed and synthesized for therapeutic evaluations since its discovery in 1988 [12–20]. However, the structural generality, non-specificity, and preferential ligand binding of the

![Figure 1: ETs; (a) ET-I, (b) ET-II, (c) ET-III, and (d) SRT-6b.](image-url)
endothelin receptors, *i.e.*, ET$_A$ and ET$_B$ types, cross-talk, common/multiple-bindings, and cross-reactivity, seemingly, subdued the progress in the ET-based therapy developments. Interestingly, the endothelin receptor types (ET$_A$, ET$_B$, and ET$_{III}$) mediate ETs regulatory actions, and the ET-I, being the biologically most potent and predominant member of the endothelin peptide family, preferentially binds to ET$_A$ receptors, whereas the ET$_B$ type receptors entangle both the ET-I and ET-III peptides [21]. The ET$_B$ receptor type/ET-III axis is also considered as an endogenous “antagonist” system of the body opposing the effects of ET$_A$/ET-I mediated bioactivities. The ET$_B$ receptor types mediate both the vasoconstriction and vasodilation activities in different tissues and organs, and seem to be have an elaborate, though probably non-specific, restricting control mechanism. Therefore, the selective ET$_A$ antagonist development for therapeutic applications have been preferentially taken up from the drug discovery viewpoint, involving the non-peptidic or mixed peptidic structural components, pharmacophore mapping, peptidomic variants, and alternates of known antagonists more frequently, and more recently [22–26].

The ET’s ET$_A$-receptor has been found to be responsible for growth promotion and pro-inflammatory activities which are responsible for developing chronic diseases, *i.e.*, atherosclerosis, hypertension, renal malfunctions, and heart failure, primarily owing to the elevated ET-I levels [27,28]. The ETs, which are potent vasoconstrictor peptides, are also considered to be associated with several other diseases of lungs, kidneys, vascular system, reproductive organs, nervous system, heart failure, and cancers. These ETs are expressed ubiquitously by the stress-responsive regulators with both beneficial and detrimental effects, specifically with the ET-II and ET-III isoforms [29–32]. The diseases associated with cell growth and inflammatory activation, *i.e.*, arterial hypertension, glomerulo-sclerosis, and immune mediated malfunctions involving cancers, connective tissues ailments, and chronic allograft rejections, as well as metabolic diseases of obesity and diabetes, are turning to be renewed challenges for developing endothelin-antagonist-based therapies [33,34].

Several studies have thus led to the discovery of selective ET$_A$ receptor antagonists (including non-selective ET$_A$ and ET$_B$ antagonists) and pharmacophore optimization for the purpose [35]. The preclinical and clinical studies have clearly established that these antagonists are effective in treatment of essential hypertension, pulmonary hypertension, heart failure, and atherosclerosis [36–40]. Advances in this area resulted in the US-FDA approval of the first orally active antagonist Bosentan followed by Ambrisentan for pulmonary hypertension and vasoconstriction [41,42]. However, the role of anti-endothelin therapy in the treatment of cardiovascular and other diseases, and determinations of the roles of the selective receptor antagonism vs mixed ET$_{A/B}$-receptor(s) antagonism in human diseases, certainly needs further elaboration from the structure activity relationship and structure property relationship (SAR and SPR) viewpoints.

The SAR studies on the structural requirements of ET-I to exhibit pressor and depressor responses are available. The oxidation of the methionine amino acid at position-7, Met$^7$, resulted in retained hypotensive activity of the analog, while the removal of the natural Cys$^1$–Cys$^{15}$ disulfide bridge led to weak agonistic action with both the pressor and depressor types’ activities of the ET-I analog. The formylation of terminal Trp$^{21}$ amino acid resulted in complete loss of the activity. These results indicated that Met$^7$ and the indole moiety of the Trp$^{21}$ are important structural requirements for the expression of activity of ET-I, whereas the intramolecular loop structure, the cyclized part, is less important. The study also provided further evidence that the depressor and pressor effects of the ET-I are mediated through different receptors [43]. The Asp$^{18}$ and Ile$^{19}$ AA residues replaced by alkyl spacers of various (chain) lengths of the C-terminal analog, Ph–Ph–CH$_2$–O–N–CH–CO–Phe–Asp–Ile–Ile–Trp–OH, proved the role of the central portion of the ET-I sequence, and helped to identify the N- and C-terminals as pharmacophoric regions of the ET-I sequence. The side-chains of the centrally located peptide have been shown to be irrelevant for the binding of the molecule to the receptor but the distance between the two postulated sites for the interaction of the ligand with the ET$_B$ receptor have been found to be of fundamental nature for bindings, and consequently, activity exhibitions [44].

These preliminary studies have opened up the ETs sequences for more scrutiny and newer developments in understanding of the SARs of the peptidic and non-peptidic analogs by various groups. Among the non-peptidic analogs, the sulfonamides, biphenyl sulfonamides [45,46], 4-sulfonamido-pyrimidines [47], azoles [48–50], and indan derivatives, as endothelin antagonists [51], including atrasentan (ABT-627)-based pyrrolidine-3-carboxylic acids[12], have been designed, prepared, and bioactivity evaluated. Among the peptide analogs, the tripeptide antagonists with unnatural AA have showed that the first AA, in the sequence, need to have a high dependency for hydrophobic residues, while the second AA can be aromatic hydrophobic AA, and the third AA must be a D-isomer as part of the minimum pharmacophore requirements based on the stereo-electronic and conformational dependency [52]. The observation that the sequence heterogeneity at the N-terminal portion of
AA from positions 4–7 showed marked biological activity differences between ETs, and SRT-6b further confirmed the role, outreach, position, and spacing importance of each of these AA. The fact that the vasoconstrictor activities of the ET-II, ET-III, and SRT-6b have been found to be one-half, one-60th, and one-third of that of the ET-I, respectively, the structural heterogeneity as well as sequence isosteric nature and homogeneity/similarities, which are well pronounced and confirmed with defined and definite levels of biological activity of the ET-I, ET-II, ET-III, and SRT-6b analogs, are proof of the concept [53]. The fact that the physico-chemical properties and molecular attributes of these molecules affected their receptor selectivity, receptor binding extents, and affinity, and the biological activity exhibitions have made them a prime target for SAR and SPR studies [54]. Moreover, a monocyclic ET-I analog, devoid of Cys² and Cys¹¹ AA which were replaced by Ala and Ala AA, had showed one-third of the activity of the ET-I, while the diaminocarbonyl analog turned out to be nearly inactive, thereby indicating the role of the termini groups. However, no significant changes were observed in major bioactivities of the isoform ET-I with replacements of Ser⁸, Ser⁹, Leu⁶, Met⁷, Lys⁹, Tyr¹³; Trp²¹ by Ala, Ala, Gly, Met(0), Leu, Phe; and Tyr or Phe, respectively. On the other hand, the replacement of Asp⁸, Glu¹⁰, and Phe¹⁴ by Asn, Gln, and Ala, respectively, resulted in complete loss of the biological activity. These results indicated that the two disulfide bonds in ET molecule are not essential for the expression of vasoconstricting activity, but both the terminals’ amino and carboxyl groups, carboxyl groups of Asp⁸ and Glu¹⁰, and the aromatic group of Phe¹⁴ seemed to be contributing to the receptor binding and expression of the biological activity, and thus, seemed to be essential for the activity manifestation [55]. Yet, in another approach, tripeptides composed of unnatural AA, namely, linear peptide derivative, BQ-485, perhydro-azepin-1-yl-l-leucyl-d-tryptophanyl-d-tryptophan resulted in potent anti-contracting activity. The HIM-CO–Leu-d-Trp-d-Phe-(–R)-OH compounds as ET₄ receptor antagonists provided a tool for the development of therapeutic agents in the treatment of ET-I manifested disorders [56]. The study by Kimura et al. [57] established the importance of the C-terminal AA, wherein replacement of Trp²¹ with d-Trp, reduction and carbamido methylation of the four Cys residues, and cleavage at Lys⁹ significantly lowered the vasoconstriction activity of the ET-I analogs. Further, studies on hexapeptide derivatives [58], linear tripeptide derivatives, R₇R₇NC(O)–Leu-d-Trp-β-Ala-OH, derived from endothelin antagonistic cyclic-pentapeptide, BQ-123 [59], peptidomimetic analog containing p-Cl–Phe [60], and combinatorial synthesis for the optimization of the potent ET-I analogs [61], are available [62]. The ETB receptor antagonists, BQ-788 [N-cis-2, 6-dimethylpiperidinocarbonyl-1-y-methylleucyl-d-1-methoxy carbonyltryptophanyl-d-norleucine] [63] further provided the required insight into the structural requirements for the bioactivity generation, the details of which could be utilized in understanding the roles of the ETs in physiological and pathological processes.

The current study focuses on a combination of approach dealing with the SPRs of various ETs isoforms, including the effects of naturally mutated AA-based analogs, miniaturized ET analogs in ET-I, as an example applicable to, in approach, to ET-II and ET-III, since ET-I is the most potent isoform. These isoforms were studied for their molecular properties, and molecular size due to the different sequence patterns. The sequences were downsized to new structures based on the ET-I together with the SRT-6b peptide, the snake venom toxin, which has been instrumental in providing a marked differentiation for receptor affinity, and its binding preferences to the ETs receptors. The observation by earlier researchers reported that the ETs are capable of modulating the biological action in so many of the body tissues and organs prompted to undertake the thorough and dissecting analysis of the sequences of ET-I isoform, and predict their effects on the possible SPRs, which also need to be validated through appropriate receptor-binding and in vivo experimentations, in addition to the in silico experimentations and prediction studies. However, the current study focuses on the computational modeling, molecular properties and QSAR value estimations in relation to the designed sub-peptidic structures. The study also proposes the nano-scale vesiculation process as part of the plausible mechanism and role of several factors of extracellular surroundings, media, pH, structural specifications, etc.

2 Materials and methods

The molecular modeling was performed on Hyper-Chem v 7.5 (Hypercube Inc., Gainesville, FL, USA), and Advanced Chemistry Development (ACD) Freeware, Ontario, Canada. The in silico generated peptide models obtained from ET-I, II, III, and SRT-6b peptide sequence structures’ manipulation, shortening, and dissection were designed. A set of approaches were consulted [29,41,64–68]. The exercise provided different cyclic peptides of varying length where the cyclizations between the Cys¹, Cys³, Cys¹¹, and Cys¹² have been made with several different combinations of the
cyclization between these Cys residues, while the ET's tail sequence was utilized as addendum to these designed cyclic structures to provide models of different sequences. All the structures were predicted for their calculated molecular and QSAR properties, and the comparisons of the properties and molecular attributes were made. The conformation linked energy minimizations through "steep descent algorithm" in Hyper-Chem v 7.5 was performed. The most stable structure arrangement and the molecular geometries were predicted, which were sorted out for further analyses of physico-chemical characteristics and QSAR properties relationship. In order to get an empirical idea, and find rationalizations about the energy levels and behavior of the different peptide fragment models, molecular energy was minimized to get the stable and minimum energy conformation thereof. The most energetically favorable models were obtained and studied. On the outset, the minimum energy conformations of various fragments were compared to know the extent of changes in the physico-chemical properties of the fragments. All the molecules and their start set-up were simulated at 0.01 Å/cycle through 1,000 cycle's calculations. The predictions for molecular properties were estimated (atoms exhibiting non-parametric characters were opted to be ignored in the software based calculations), and molecular volume, total charge density, and electrostatic potentials of the fragments, their log P, and hydration energies were calculated.

3 Results and discussion

3.1 Optimization of ETs, SRT-6, and sequence fragmentations: Approach and design

The ETs, naturally encountered as ET-I, ET-II, and ET-III isoforms, with differentiating sequences were primary structures to begin with. Interestingly, all types of endothelin's tails have major sequence similarities with the snake venom toxin, SRT-6b sequence, but lacked the commonality of the biological activity with either being inactive, strongly active, diminished, or different activities. The study pursued to generate and rationalize the design of new structures through permutation and combination approach of the AA sequence to reach the best and optimized structure for further development. The structural similarities and differentiating points were considered in the design.

3.2 Tail and cyclic portions' analysis of the ETs and SRT-6b: Activity and property implications

For all the ETs, ET-I, ET-II, and ET-III, the hexapeptide tail sequence of AAs located from position 16–21 (single letter code sequence; HLDIIW) is similar, and is suggested to be structurally insignificant in terms of demarcation in receptor binding for biological activity elicitation; although, it has been suggested that the C-terminal hexapeptide chain discriminates between the different endothelin receptors [69]. Since this tail is similar in sequence structure for all the endothelin sub-types, the possibility for it to be distinguishing between receptor sub-types is remote. Any possibility of distinction between the receptor sub-types by the tail lies in the tail adopting different conformations and being distinctively different in the surroundings for differentiating the various receptor domains with the stereo-orientation assisted and modified by the (nearly common) cyclic part of all the ETs' sub-structures. Hence, the cyclic part of the structure may also play an important role in distinguishing the different receptor sub-types, and there is strong probability of it to do the major part in this exercise. The major AA-based structural differences of the main body cyclic part of the ETs is in the AA sequence for positions 2, 4, 5, 6, and 7. The differing AAs are Leu and Met at positions 6 and 7 of ET-I, which are replaced by Trp and Leu AA-residues, respectively, at position 6 and 7 for the ET-II, while the ET-III is completely different by the presence of Thr, Phe, Thr, Tyr, and Lys at 2, 4, 5, 6, and 7 positions of the AA sequence of its structure. However, the snake venom toxin, SRT-6b, also a 21 AA hexapeptide chain, have a different AA sequence structure as compared to that of the ETs tails (AA 16–21). For the cyclic structure part, few common AAs are available in the sequence for SRT-6b, otherwise it is largely different in its structure. This is an intriguing structure similarity/dissimilarity, seemingly, responsible for venom’s extreme cardiovascular inhibitory activity [70–72].

3.3 Peptide foldings, compartmentalization, and mutated AA: Geometric views

The main body cyclic sub-structure of SRT-6b differs from ET-I by AAs located at positions 4–7 and 12, and replaced AAs are identified as Lys, Asp, Met, Thr, and Leu. The Leu at position 12 in SRT-6b is occupied by Val in the ETs'
sequences. The SRT-6b is also known to interact and involve in competitive binding with ETs [73]. Therefore, the point of interest for the AA sequence in SRT-6b is the tail composition which is, supposedly, also responsible for the demarcation between endothelin receptor types in ETs structures. The AA positions at 17 and 19 in all ETs are filled by the Leu and Ile AA residues, while the same slot is occupied by Gln and Val in SRT-6b molecular compositions. However, with the presence of different receptors in ileum and cerebellum [74], this observation further confirmed the presence of different ET types on structures’ basis. It may also be considered that different structural domains formed from the molecular interaction and aggregation of the structures in the surrounding areas give rise to distinct compartmentalization, in their interaction area, when interacting with outside moieties, especially, water and other smaller entities, e.g., ions and atoms in the structures, pH, media polarity, surface, and molecular entities’ interactions of hydrophilic and hydrophobic nature. The axis-based alignments in the molecule in response to the external and internal factors and interactions help to formulate the compartment (Figure 2). The compartmentalization seemingly is facilitating the approach to binding, and enhancing the competitive binding by providing a different look in peptidic folding and its conformations to the ETs and SRT-6b receptors.

3.4 Molecular properties and QSAR attributes: SPR and SAR

The presence and interactions of singular molecular water, and water as part of the cluster, ions, and other small molecules at certain specified locations in the receptor area, along with the participation of inorganic, and/or other non-proteinaceous organic ligands, as well as the receptors maneuvering the binding feasibility even by difference of a single AA in the structures of SRT-6b and ETs may be very crucial. The physicochemical factors are vital and play significant roles at receptors’ interaction levels. The hydrophilicity, hydrogen binding, hydrophobicity, and electronegative characteristics of the atoms, total charge density and its distribution on the molecule, as well as the

![Figure 2: Peptide folds, ET-I based with compartmentalization, and the sequence’s geometries: Emerging structural orientations with resultant geometry and axial alignments of the molecule providing different views of the geometry, with all the back-bone atoms in the models appearing as green atoms; (a) molecule leading to compartmentalization, shown with the explicit hydrogen atoms; (b) nearly three distinct major regions are appearing; (c) primary X-axis alignment; (d) Y-axis alignment; (e) primary Z-axis alignment, probably for passage through channels; (f) secondary Y-axis alignment; (g) secondary Z axis; (h) tertiary X (as major axis), primary Y axis orientation. Structures are shown without explicit hydrogens, the space-filling models’ explicit hydrogens are white, the carbon atom is magenta colored, oxygen is red, nitrogen blue, and sulfur is yellow colored, the color scheme follows the current and all the subsequent models, unless otherwise stated.](image-url)
electrostatic potential on structural parts and fragments, are some of the parameters to look for. The paracohor, index of refraction, surface tension, density, and ring double bond equivalent (RBDE), of which the later is representative of the levels of unsaturation in the molecule and their fragments, and perhaps is being considered in the software pediction to indicate for the hydrophobic character contribution, which is prominent in receptor binding interactions. These predictors are also important to look for from different perspectives of the molecular properties of the structures. The QSAR molecular attributes predicting polarizibility, refractivity, log P, hydration energy, and molecular volume are other factors suggestively forming the SAR/SPR of different ET types, and the receptor demarcation activity by these ET motifs is pursued by the molecular properties in conjunction with the QSAR predictors. The prepared monocyclic fragment analogs of ET-I for exploring the importance of the bicyclic structure of ET-I to its predictive molecular and electronic properties showed low micro to high nano-molar binding affinities, and were functional antagonists of ET-I for induced accumulation of inositol phosphates. However, one analog possessed mixed antagonist/agonist activity at the two ET receptor subtypes [75–77]. The current work involved the in silico modeling and generated analogs of the ETs. The mapping of ET-I showed the 2D contour graph for the total charge density and electrostatic potential localized in the cyclic structure part of the molecule, which apparently was suggested as inactive, while the common to all ET was the tail sequence (AA 16–21). Hence, from the viewpoint of design and in silico study, the cyclic structure part needed more elaborate understanding to distinguish between the receptor sub-types, which is evident from their differing bioactivity and receptors’ affinities. On comparing the molecular attributes and QSAR data (Table 1) for ETs and SRT-6b, the log P and hydration energy differences are remarkable, while other characteristics and QSAR properties also differ, interdependent of their structures. The structures of the ET-I, II, III, and SRT-6b are represented in Figure 3. The ET-I molecular modeling based electrostatic potential was predicted high, mainly localized in the loops (S–S bonds and non-tail AA 16–21 area of the structure), while the total charge density was moderate and distributed in the similar regions as that of the ET-I’s electrostatic potential. The ET-II exhibited high electrostatic potential and high total charge density distributions, both being more than the ET-I, while the ET-III showed lesser electrostatic potential and total charge density than the ET-II but more than the ET-I. The SRT-6b exhibited these molecular characteristics nearly equal to ET-III. Of the ETs, the ET-I being most potent of the ETs, the high electrostatic potential and medium levels of charge density seemed to be favored for ET-I functions, and an antagonist activity elicitation is expected to be triggered through the competitive binding.

### 3.5 Role of Cys residues, sequence shortening, and miniaturized ET: Dissection of the big ET

The big ET sequence of 38 AA was miniaturized to smaller 21 AA sequence to render the required biological activity. The elicitation is naturally possible after cleavage of the Trp–Val (21–22 AAs) bond by protease enzyme, and the biological activity is dependent on the cleavage process, and proceeds as the intermediate action, which is facilitated physico-chemically. Interestingly, the 22–38 AA sequence for human ET-I contains no Cys residue. The small ETs sequence stabilize them by attaching to other ligand through physical interactions, which provides physiochemical changes necessary to elicit the biological response when interacting with the receptor. However, it was also decided to take an approach to find the minimum sequence possible for the ET-I on the 21 AA sequence, and permute and combine sequences for finding a smaller possible sequence to entail the 16–21 AA residues to it, and produce a mini ET sequence equivalent to ET-I as a template by inducing the sequence optimization as described in structure dissecting approach. The energy diagram, electrostatic potential, and other comparative molecular attributes were also predicted.

### Table 1: Molecular attributes and QSAR properties of the ETs and SRT-6b*

| Property          | ET-I   | ET-II  | ET-III | SRT-6b |
|-------------------|--------|--------|--------|--------|
| Molar refractivity| 629.50 | 648.98 | 676.06 | 641.30 |
| Molar volume      | 1,957.0| 1,979.7| 2,056.30| 1,974.80|
| Log P             | 4.98   | 7.95   | 9.03   | 5.40   |
| Hydration energy  | −40.20 | −47.61 | −57.28 | −57.79 |
| Paracohor         | 5,227.60| 5,335.2| 5,564.16| 5,339.66|
| Index of refraction| 1.556 | 1.569  | 1.571  | 1.562  |
| Surface tension   | 50.9   | 52.7   | 53.6   | 53.43  |
| Density           | 1.273  | 1.286  | 1.285  | 1.298  |
| Polarizability    | 249.55 | 257.27 | 268.01 | 254.23 |
| RDBE              | 43     | 49     | 51     | 45     |

* Molar refractivity expressed as Å³ (cubic Angström), volume as Å³, hydration energy in kcal/mol, density in g/cm³, surface tension in dyne/cm, paracohor as Å³, polarizability as Å³; insignificant variations have been omitted. The log P and hydration energy were predicted from Hyper-Chem 7.5, and rest of the properties were calculated from the ACD software.
3.6 Cavity modeling, Cys–Cys bonds, hydrogen bondings: Evolution of electronically active part

The backbone comprising the α-carbons of the AAs and the amide bonds were modeled for their minimum energy conformation (MEC) energy conformations based geometric orientations (Figure 4). All the 21 AAs produced a tailed cavity and AAs 1–15 were part of the cavity (Figure 4c), while the tail sequence AA 16–21 were not in the cavitation, and the atoms on the backbone are visualized in Figure 5. The structural stacking (overlaps) and geometric changes were facilitated by the energy requirements, and the Cys–Cys bondings of the ET-I molecule (Figure 6), hydrogen bonding effects, and internal axes from 1–2 and 3–4 showing the atoms and the residues distribution ratio in the ET-I molecule along the backbone (shown as broken lines other than the hydrogen-bondings),
Figure 3: Structures of ETs and SRT-6b; (a) ET-I and (b) ET-I space-filling model; (c) ET-II and (d) ET-II space-filling model; (e) ET-III and (f) ET-III space-filling model; (g) SRT-6b and (h) ET-III space-filling model; all structures and models are minimum energy conformers. Structures are shown without explicit hydrogens, the space-filling models' explicit hydrogens are white.
played a part in the compartmentalization. The distribution along the axes 1–2 and 3–4 indicated that the ET-I tail (AA 16–21) together with AAs positioned at 13,14, and 15 forms the distinct distribution part as equal weightage with the counterpart of the distribution, the two Cys–Cys cyclizations (AA 1–15 and 3–11) and the residue 12 (Figure 7). The exclusivity of the chain is also exhibited in the electrostatic potential and total charge distribution (Figure 8), which is localized in the non-tail part of the ET-I structure. This also may help to refute the notion of the cyclic part (AA 1–15) being inactive as this part has more pronounced electron density, charge distribution, and consequently prone to chemical/biochemical interaction with energy requirements to be active.
3.7 Prototypical template, sequence analysis, and replaceable tail: Generation of new chemical entity

The ETs without tails are a bicyclic structure identified as cyclic structures, A and B, with S–S bonds between two Cys units at positions 3 and 11, and at positions 1 and 15, together with the interconnecting atoms and bonds of the peptidic structure between them. One cyclic unit contains 8 AAs identified as sequence Cys3–Ser–Val–Tyr–Phe–Cys15 with S–S bonds between AA 1 and 15, and AA 3 and 11. The other unit constituted of 9 AAs identified as a sequence Cys3–Ser–Leu–Met–Asp–Lys–Glu–Cys11 with S–S bonds between 3 and 11. The two structures are deduced after Cys3 and Cys11 positions taken as common between both the cyclic parts. The addition of tail from either ETs, or from the SRT-6b provided two mini ET prototypical templates. Another combination making an interesting structural subtype for the monocyclic mini ET ring is produced by the modeling of ET-I, which constituted 12 AAs identified in sequence as Cys3–Cys11–Val12–Tyr13–Phe14–Cys15–His16–Leu17–Asp18–Ile19–Ile20–Trp21 incorporating the hexapeptide tail with AAs from 16–21 with Cys–Cys S–S bond between AA at positions 3 and 11. In an attempt to partially open the ET-I
structure, the AA linkages between AAs at positions 3 and 4 were cleaved to give rise to two tail like structures linked to AAs 11 and 15.

The original hexapeptide side chain was intact in its attachment to the Cys\(^{15}\) AA as viewed on the ET-I models. The additional side chain thus generated contained AA Ser\(^{4}\) to Glu\(^{10}\) linked sequentially, and joined to the monocyclic structure at Cys\(^{15}\). Thus, the monocyclic ring comprised of AA Cys\(^{1–2}\)–Ser\(^{3}\)–Cys\(^{5–11}\)–Val\(^{12}\)–Tyr\(^{13}\)–Phe\(^{14}\)–Cys\(^{15}\). The shortening of the cyclic sub-structure contained the AA identified in the backbone of ET-I, where side chain is tail-end in both the cases. However, an analysis on the majority of the occurrences of AA in ET-I revealed that the Cys (at 4 positions, i.e., 1, 3, 11, and 15) is the most common AA followed by the Ser(3), Leu(2), Asp(2), and Ile(2) where Ile, found 2 times, is specific to the hexapeptide tail chain. The sequence derived could be Cys–Ser–Leu–Asp–Cys conjoined by S–S bond between first and last Cys residues. A 3D analysis of ET-I showed the concentration of aromatic ring containing AA in one domain probably facilitating the binding [78]. The sequence can be modified to include the hexapeptide chain with the most common AA sequence Cys–Ser–Leu–Asp–Cys and with the sequence Cys–Ser–Leu–Asp–Cys–Ile, although the change in hexapeptide constitution is not favorable and is not recommended keeping in view of the homology with other ETs tails (AA 16–21), i.e., ET-II, and ET-III.

The differential biological activity of ET-II and ET-III are different and their receptor affinity separate with ET-II have certain cross-affinity. The fact that the ET-II has only two different AAs (Trp at 6, and Leu at position 7) as compared with the ET-I, while the ET-III have a total of 6 different AAs, of which all the 6 are of the non-tail part, and localized in the cyclic structure part of the ET-III, as compared to the ET-I. The AA differentiation has contributory roles in the receptor affinity of the three distinct ETs, through the interactions at the molecular and atomic levels. The AA occurrence criteria and repeating sequence observations for these AA provided Cys–Ser–Trp–Leu–Asp–Cys as the other template for ET-II with the common ETs hexapeptide side chain (AA 16–21) attached. The Cys–Ser–Trp–Leu–Asp–Cys–Ile is the other sequence analyzed for its analogy for the ET-II sequence shortening as a ligand for probably being binding facilitator. The ET-III analysis produced the AA sequences Cys–Thr–Phe–Tyr–Lys–Cys, Cys–Thr–Phe–Tyr–Lys–Asp–Cys and Cys–Thr–Phe–Tyr–Lys–Asp–Cys–Ile with the common hexapeptide chain where both the Cys\(^{2}\) are joined by S–S bonds to give the monocyclic structures. The energy profile and conformational analysis put the sequences in similar category of physico-chemical parameters. The AAs Asp and Ile in the proposed common AA-based sequences can be deleted in lieu of the hexapeptide chain attachment. The extra Cys was provided for the S–S linkage. The SRT-6b hexapeptide tail is also an important sequence moiety needing evaluation in the changed structural domains, and in the sequence shortening exercise with the detailed biological evaluation for these sequences is necessary in future.

### 3.8 Cys AA combinations and further shortening of the ETs: New and shortened sequences

The Cys combinations can be tried for further shortening of the long sequences of the ETs. The first approach was keeping the Cys residues as Cys\(^{1–2}\)–Cys\(^{3}\)–Cys\(^{11}\)–Cys\(^{15}\) with S–S bonds between Cys\(^{1–2}\)–Cys\(^{15}\) and Cys\(^{1–2}\)–Cys\(^{11}\) with the hexapeptide tail from the ETs. The further shortening can give Cys–Ser–Cys with S–S bonds between both the Cys residues, and any one of the Cys residues attaching the common hexapeptide side chain from the ETs. The sequence for ET-I and SRT-6b based moiety will be of interest to compare as there exists lesser commonality between them. The ET-III have different AA sequences than the ET-I, whereas the ET-III differs in a total of six AAs and all of them are in the non-tail part of the structure. The ET-III is also considerably different in its AA constituents as compared to the ET-II molecule. The structural differences speak for the varying molecular properties, QSAR values (Table 1), and consequently the differences in the receptor type affinity. The ET-III was distinctly different than the ET-I, and seemingly was more close to the SRT-6B structure. The AA commonality and differences, and their respective physico-chemical evaluation, and biological activity response profile as well as the effects of the shortened sequences are worth deliberating. The ET-I sequence in another combination approach provided Cys\(^{3}\)–Cys\(^{11}\)–Val\(^{12}\)–Tyr\(^{13}\)–Phe\(^{14}\)–Cys\(^{15}\) and Cys\(^{3}\)–Val\(^{12}\)–Tyr\(^{13}\)–Phe\(^{14}\)–Cys\(^{15}\) with S–S bonds between Cys residues in analogy to the ET-I (Cys\(^{3}\)–Cys\(^{15}\), Cys\(^{3}\)–Cys\(^{15}\)), between Cys\(^{3}\)–Cys\(^{15}\), between Cys\(^{3}\)–Cys\(^{15}\), and between Cys\(^{3}\)–Cys\(^{11}\) as derived from the ET-I sequence with the hexapeptide attached. The changeover of the S–S bond relationship in ET-I to Cys\(^{3}\)–Cys\(^{11}\) and Cys\(^{3}\)–Cys\(^{15}\) provided the sequence as Cys–Cys–hexapeptide tail, and Cys–Ser–Cys–hexapeptide tail with feasible S–S bonds between the Cys residues. The non-hydrophilic AA in lieu of Cys residue in S–S linkage cross-over model can also be tried. The AAs can be Ala, Val, Leu, and Pro.
The cyclic part of the ET-I will give the Cys–Ser–Val–Tyr–Phe–Cys, Ser–Cys–Val–Tyr–Phe–Cys, Cys–Tyr–Phe–Cys from the substructure Cys$^1$–Ser$^2$–Cys$^3$–Cys$^{11}$–Val$^{12}$–Tyr$^{13}$–Phe$^{14}$–Cys$^{15}$ and Cys–Ser–Leu–Met–Asp–Lys–Glu–Cys, Ser–Leu–Met–Asp–Cys–Glu with S–S bonds between terminal Cys, and amide linkage in last example between Cys and Glu from the ET-I substructure Cys$^3$–Ser$^4$–Cys$^5$–Leu$^6$–Met$^7$–Asp$^8$–Lys$^9$–Glu$^{10}$–Cys$^{11}$. The hexapeptide chain from ETs and SRT-6b as tail attachments at the carboxyl and N-terminal have also been analyzed for their molecular attributes and conformational analysis. The dissecting approach to ET-I producing various shorter peptides is depicted in Figure 9. The generated structures, S1–S18, were estimated for their molecular and QSAR properties (Table 2). The electrostatic potential and charge density were observed as high, moderate, and low in nature based on their intensity expressed as eccentric lines in high, low, and medium densities with localizations/distributions on the structures. Among the high exhibits of the electrostatic potentials were the structures S1, S4, S5, S8, S9, S12, S13, S14, S15, S16, and S17, Bosentan, Mecitentan, and other antagonists, e.g., BQ-123, 485, and 788, while the medium

![Figure 9: ET-I: structure dissecting approach, all AAs represented by a number are continuous natural chain of ET-I unless stated in the figure, all Cys–Cys S–S bonds are straight lines unless otherwise stated; for figures S16, and S17, the broken lines represent the S–S bonds between the two Cys residues, Cys$^1$–Cys$^{15}$.](image-url)
Table 2: Molecular attributes and QSAR properties of the ETs and SRT-6b derived structures (S1–S18)

| Molecular properties | S1      | S2      | S3      | S4      | S5      |
|----------------------|---------|---------|---------|---------|---------|
| Molar refractivity   | 442.55  | 447.39  | 431.31  | 251.82  | 458.17  |
| Molar volume         | 1,378.5 | 1,195.65| 1,247.20| 677.65  | 1,243.95|
| Log P                | 1.14    | 0.71    | 0.28    | −1.49   | 3.46    |
| Hydration energy     | −38.37  | −39.87  | −44.08  | −17.37  | −31.34  |
| Parachor             | 3,637.90| 3,668.60| 3,550.14| 2,036.50| 3,733.86|
| Index of refraction  | 1.55    | 1.67    | 1.608   | 1.665   | 1.6580  |
| Surface tension      | 48.40   | 88.65   | 65.63   | 81.55   | 81.15   |
| Density              | 1.283   | 1.48    | 1.370   | 1.45    | 1.41    |
| Polarizability       | 175.44  | 177.36  | 170.98  | 99.83   | 181.63  |
| RBDE                 | 33      | 32      | 31      | 17      | 33      |

| Molecular properties | S6      | S7      | S8      | S9      | S10     |
|----------------------|---------|---------|---------|---------|---------|
| Molar refractivity   | 606.76  | 468.86  | 371.14  | 287.85  | 494.28  |
| Molar volume         | 1,819.20| 1,428.60| 1,098.80| 862.10  | 1,463.30|
| Log P                | 7.72    | 6.12    | 5.84    | 1.06    | 4.02    |
| Hydration energy     | −48.67  | 14.51   | −27.06  | −19.09  | −44.63  |
| Parachor             | 4,992.66| 3,892.26| 3,019.96| 2,460.66| 4,157.90|
| Index of refraction  | 1.581   | 1.570   | 1.590   | 1.582   | 1.590   |
| Surface tension      | 56.70   | 55.03   | 57.00   | 66.33   | 65.13   |
| Density              | 1.296   | 1.292   | 1.282   | 1.395   | 1.353   |
| Polarizability       | 240.54  | 185.87  | 147.13  | 114.11  | 195.95  |
| RBDE                 | 46      | 34      | 30      | 14      | 30      |

| Molecular properties | S11     | S12     | S13     | S14     | S15     |
|----------------------|---------|---------|---------|---------|---------|
| Molar refractivity   | 425.75  | 632.19  | 426.75  | 633.10  | 625.33  |
| Molar volume         | 1,252.7 | 1,853.90| 1,152.55| 1,751.35| 2,000.50|
| Log P                | 0.67    | 5.66    | 0.02    | −3.31   | −2.27   |
| Hydration energy     | −45.12  | −49.51  | −44.18  | −46.28  | −36.44  |
| Parachor             | 3,561.06| 5,258.34| 3,530.46| 5,227.66| 5,179.76|
| Index of refraction  | 1.595   | 1.597   | 1.662   | 1.659   | 1.537   |
| Surface tension      | 65.20   | 64.70   | 88.05   | 86.25   | 44.90   |
| Density              | 1.369   | 1.345   | 1.48    | 1.45    | 1.236   |
| Polarizability       | 168.78  | 250.62  | 169.18  | 250.98  | 247.90  |
| RBDE                 | 26      | 42      | 27      | 43      | 44      |

| Molecular properties | S16     | S17     | S18     | ET-Tail (AA^{16–21}) | SRT-6b-Tail (AA^{16–21}) |
|----------------------|---------|---------|---------|-----------------------|--------------------------|
| Molar refractivity   | 423.06  | 629.50  | 423.06  | 209.95                | 204.29                   |
| Molar volume         | 1,355.8 | 1,957.0 | 1,355.8 | 622.13                | 589.53                   |
| Log P                | −1.09   | 3.86    | 5.76    | 4.54                  | 1.72                     |
| Hydration energy     | −39.95  | −37.50  | −56.19  | −14.46                | −19.11                   |
| Parachor             | 3,530.40| 5,227.6 | 3,530.4 | 1,721.7               | 1,678.94                 |
| Index of refraction  | 1.536   | 1.556   | 1.536   | 1.590                 | 1.609 2                  |
| Surface tension      | 45.90   | 50.9    | 45.9    | 58.60                 | 65.73                    |
| Density              | 1.264   | 1.273   | 1.264   | 1.279                 | 1.351                    |
| Polarizability       | 167.71  | 249.55  | 167.71  | 83.23                 | 80.99                    |
| RBDE                 | 27      | 43      | 27      | 16                    | 17                       |
were S2, S6, S7, S10, S11, and S18, and the low was S3. The charge intensity and localizations were high in S1, S6, S9, S10, S12, and S14, and medium levels were exhibited by S2, S4, S15, and S18, and Bosentan, Mecitentan, and other antagonists, while the low levels of the charge density were found in the structures S2, S3, S5, S7, S8, S10, S11, S13, S16, and S17. A closer look favored the high electrostatic potential with low to moderate/medium levels of charge density. This favored the structures S4 and S15, while other considerations were made to include other structures too (Figure 10).

3.9 Characteristics of the tail chain (AA 16–21) of ETs and SRT-6b

The ETs common tail (Figure 11) chain comprising the hexapeptide (AA 16–21, His–Leu–Asp–Ile–Ile–Trp, HLDIIW, single letter code sequence) and the SRT-6b hexapeptide (AA 16–21, His–Gln–Asp–Val–Ile–Trp, HQDVIW) were compared for their molecular properties and the QSAR values (Table 2). The major differences in the properties were observed in the log P values (ETs chain: 4.54 and SRT-6B chain: 1.72), hydration energies (ETs chain: −1,446 kcal/mol and SRT-6b chain: −1,911 kcal/mol), and molar volume (ETs chain: 622.13 Å³ and SRT-6b chain: 589.53 Å³).

The mutation approach of replacing the AA were utilized to further dwell upon the characteristic changes in the substructures and ETs (ET-I, II, and III) were provided with the SRT-6b tails (AA 16–21, His–Gln–Asp–Val–Ile–Trp), as well as the SRT-6b was attached with the replaced tail of the ETs. The QSAR and molecular properties estimation (Table 3) showed that the ET-III and SRT-6b produced characteristics in the same range, while the ET-I and ET-II were of the same league in their molecular properties, especially the log P and hydration energy together with the refractivity.
Figure 10: ET-I derived structures; structures of: (a) S1 (AA 1–15 and 3–11 Cys–Cys S–S bonds) and (b) S1 space-filling model; unusual Cys–Cys bonds (AA 1–11 and 3–15 Cys–Cys) models; (c) S16 and (d) S16 space-filling model; (e) S17 (AA 1–11 and 3–15 Cys–Cys S–S bonds) and (f) S17 space-filling model; (g) S18 (AA 1–15 and 3–11 Cys–Cys S–S bonds and the Tail chain AA 16–21 is attached as reversed sequence, AA 21–16 = Tyr–Ile–Ile–Asp–Leu–His), and (h) S18 space-filling model. All structures are shown without explicit hydrogens, the models' explicit hydrogens are white.
The mutant structures have also been designed and some of the mutant points for ETs have been identified through NMR analysis and that confirms the existence of mutants [79].

### 3.10 Comparative study of the properties: Antagonists evaluation

The molecular properties and the QSAR values estimation of the model ET-I antagonists, Bosentan, Mecitentan (Figure 12), BQ-123, BQ-485, and BQ-788 were carried out (Table 4).

The studied antagonists have a range of \( \log P \) values, hydration energies from lowest (−6.38 kcal/mol) to highest (0.28 kcal/mol) and differing molecular sizes. The values have importance in their range which entails that the very high molecular and QSAR property values are for the high molecular weight (MW) products, and that the low MW products designed on the ETs structural analysis can also be potent and selective based on the SPR relationship when compared with the known antagonists.

However, for the designated products and the ETs, there were ranges of \( \log P \) from 4–9 and hydration energies between 40 and 60 kcal/mol that were observed for the ETs structures. The designed chemical entities were expected to fall in this range for probable bioactivity as ET-I antagonist. Nonetheless, the \( \log P \) values and the hydration energies were in the ranges of 1–7 and −13.50–0.28 kcal/mol, respectively, for the known antagonists (Table 4). The ET-I derived

![Figure 11: Tail chain structures (AA 16–21); (a) ETs tail chain structure (without explicithydrogens shown); (b) ETs chain structure space-filling model; (c) SRT-6b chain structure (without explicit hydrogen atoms shown); (d) SRT-6b chain structure space-filling model.](image)
and designed compounds were expected to exhibit these value ranges. Some of the designed structures, S1–S4, were near in the desired geometric shape and size range in accordance with the antagonistic molecules, while the structures S16–18 were near to the ET-I. The structures, S5–S12, were in the intermediate ranges of the size and shape as compared to the ET-I. The Q SAR and the molecular properties comparison also indicated these preferences, and the structures, S2, S4, S11, S14, S15, and S16, seemed near the mother ET, ET-I, in comparative outcome of the analysis, although the variations in the properties were pronounced as compared to the molecular shape, size, and the presence of double bonds equivalents, together with the other properties of parachor, hydration energy, polarizability, etc. (Table 2).

### 3.11 Vesiculation: A plausible approach

The functions and transport of different biomolecules may require nano-vesiculation that is considered safe for transport, stability, and supposedly works as dynamic entity to perform physiological functions of inherent and dictated nature at the transported site. Protection against cytosolic calcium ions overload in cellular stress conditions and cell injury are also known to be maneuvered through vesiculation at nanoscale of the membranes. A rapid vesiculation of peptidic molecules have also been suggested. The surroundings, structural, and physico-chemical properties led compartmentalization due to selective constrictions in the structural network producing reduced volume of the structure may further tend to minimize size and molecular weight and geometry characteristics leading towards generation of vesiculation. The structural compartmentalization, compartments of geo-spatial changes, seemed to be an outcome of several factors, including asymmetric presence of calcium ion [80–83]. The interconnected open structural parts formed as the final, or intermediate shape as tubular entities, which enables the free diffusion of proteins and calcium ions [84] can be a format for structural change. Nonetheless, the structural integrity and intrinsic characteristics to retain and preserve the primary structure puts the peptidic

### Table 4: Molecular and QSAR properties of the known ET-I antagonists

| Molecular properties    | Bosentan | Macitentan | BQ-123 | BQ-485 | BQ-788 |
|-------------------------|---------|------------|--------|--------|--------|
| Molar refractivity      | 143.69  | 125.55     | 155.26 | 173.91 | 165.21 |
| Molar volume            | 416.0   | 351.2      | 618.79 | 592.10 | 762.95 |
| Log \( P \)             | 1.06    | 3.21       | 4.67   | 3.26   | 6.96   |
| Hydration energy        | –13.49  | –2.61      | –6.38  | –4.41  | 0.28   |
| Polarizability          | 56.96   | 49.77      | 63.81  | 70.47  | 65.35  |
Figure 13: Plausible approach to nano-vesiculation; (a) surface interaction; (b) interfacial activity; (c) single-dimension interaction; (d) another dimension based interaction and structural motifs assembly; (e) increased interaction; (f) twisting of the structural motifs’ direction; (g–i) multi-directional increased movements and interactions; (j) lateral compression of the assembled structural motifs; (k) compression leading to volume reduction, and counter-compression yielding to increased interfacial interactions, and further compression and reduction of volume; (l) formation of a prototypical nano-vesicle, entrapping ions, water molecule, and small molecular weight entities.
These vesicles can be nano-sized based on the peptidic structure size and molecular weight, as well as their propensity to be nano as dictated by the molecular and physico-chemical interactions, and the role of surroundings and surface in contact. The growing nanostructures may go through the compartmentalization of the molecular area based on their physical interactions towards a tube-like structure, or a cavitation to give the supramolecular structure of assembled peptidic structures of diminished size which are prone to more intense, multi-cornered interactions and that may tend to be rounding and layering through assembly and collection of many peptidic structures with their irregular and haphazard interactions among themselves to produce an ultimate vesicular form [81]. Additionally, the factors which tend to vesiculize the peptidic structures, including surface interactions within itself and other entities, especially of lipidic nature, hydrogen-bonding and presence of hydrophobic groups and their interactions, energy requirements, initiation to geometrical prototype development and dip for curvature, compartmentalization, backbone layering of the molecular templates under the influence of physico-chemical forces, and interactions as peptide nanostructures lead through endogenously self-initiated, and self-assembled, and contributed by the thiol reactivity of proteinic cysteines are suggested [93,94]. However, the self-assembly in lipidic surroundings preferentially led to small nano-disk formation of the polypeptide molecules, and the nano-disks are thermodynamically favorable [95,96]. Experimentally, as recently reported [97,98], the surface roughness retards the molecular stretching, and geometric folding to provide diminished crystallinity for certain polymeric nanostructures, and any untoward attempt to crystallize the peptidic structure seemingly leads to vesicle formation with the growth of the nano-mass till its critical/threshold size is reached, also controlled by the factors of surface, surroundings, and intrinsic physico-chemical interactions of the developing nano-structure. The interactive, interfacial assembled peptides are prone to vesiculation which is suggested to be either generated through formation of nanodisks which comparatively are more energy preferred [89,99–101], or alternatively through different route. The protein-based structures follow the lead of the other self-assembled bio-nanostructures, e.g., DNA and RNA. The self-assembled virus complex encapsulate and transport the viral DNA for host cell delivery. The natural polypeptides of 20 AAs with differing properties, and a minimum of 4 structurally similar nucleotides have been found to introduce towards forming the nanostructures [102]. Vesicular shapes self-assembled from folded and globular protein molecules, and under aqueous media, as well as thermally-triggered self-assembled vesicles with roles from different factors, including the interfacial interactions among the polymeric structures that is strong for lipidic vesicles preparation, and the other parameters, i.e., phase transition, hydrogen bonding, hydrophilicity, electrostatic repulsion, hydrophobic interactions, changes in the surface area of the layer, and lateral pressure in initiating the vesiculation have also been suggested [103–105]. The vesiculation steps are predicted based on the physico-chemical interactions, surroundings, and resultant structural changes [86] including generation of passage for transport, or other activities. These activities may happen within seconds to provide needed passage, protection, and lead to the formation of different shapes, preferably as a vesicle, or a spherical cage (Figure 13) [87–105]. The energy in solvation, hydrophilic attractions, systemic and cellular components interactions, repetitions in structure, presence of ions, zwitterionic and dipolar nodes, and size-limitation (nano-cut) at structural/atomic scales, and the stability requirements catapult the longer biopolymeric structures to nano-shaping.

4 Conclusion and prospects

ETs and SRT-6b structural analyses generated sequences of shorter AA lengths with the molecular attributes and QSAR properties, of which some matching the original ET sequence. Certain sequences were in silico generated through different approaches of molecular analysis which showed promising molecular properties at par with the mother ET which can be developed to be utilized as ET antagonist. More advanced functional analysis of ETs structures’ through various approaches and methodology and synthesis of the ETs-derived structures awaits, which is not only for evaluating additional pharmacological studies using the specifically designed ET antagonists, but also generation of genetically altered animal models for testing, especially rodents, owing to the feasibility and observations that the majority of ET-based design and bio-testing studies have been performed on the rodent models. The bioactivity evaluation as an approach where conditional loss-of-biofunction(s) and gain-of-biofunction(s) manipulations to check the hypothesis is desired. Studies in feasibility, mechanism, process, and simulations for vesiculation of ETs, including big ET, together with their property and biofunctions need to be experimentally studied.
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