Molecular Evolution in a Peptide-vesicle System

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Abstract: Based on a new model of a possible origin of life, we establish an efficient and stable system undergoing structural reproduction, self-optimization and molecular evolution. This system is being formed under realistic conditions by the interaction of two cyclic processes, one of which offering vesicles as the structural environment, the other supplying peptides from a variety of amino acids as versatile building blocks. We demonstrate that structures growing in a combination of both cycles have the potential to support their own existence, to undergo chemical and structural evolution and to develop unpredicted functional properties. The key mechanism is the mutual stabilization of the peptides by the vesicles and of the vesicles by the peptides together with a constant production and selection of both. The development of the proposed system over time not only would represent one of the principles of life, but could also be a model for the formation of self-evolving structures ultimately leading to the first living cell. The experiment yields clear evidence on a vesicle-induced accumulation of membrane-interacting peptide which could be identified by liquid chromatography combined with high-resolution mass spectroscopy. We found that the selected peptide has an immediate effect on the vesicles, leading to i) reduced vesicle size, ii) increased vesicle membrane permeability, and iii) improved thermal vesicle stability.

Keywords: Origin of life; evolution; molecular evolution; prebiotic chemistry; peptides; vesicles

1. Introduction

It is generally accepted that complex prebiotic structures do not appear accidentally, but instead form in a long-term process facilitated by random variation, selection and reproduction. This process, representing a very general form of evolution, must have dominated even the very early steps leading to very primitive precursors of a living cell. It must have been based on the building blocks of prebiotic chemistry which potentially have been formed in many different locations on or near the early Earth [1-9].

An important fraction of prebiotic compounds are amphiphilic substances, the precursors of lipids. Their unique tendency to form various mesostructures, most prominently multilayers and
vesicles with double-layer membranes, makes them a natural starting point for cell-like compartments [10-12]. They also have the capability to select and accumulate other prebiotic molecules [13,14], especially if those molecules are amphiphilic as well [15,16]. Those amphiphilic mesostructures may occur naturally in a variety of environments, such as shore lines, hot springs [11,17] or even in bulk liquid phases.

A very special environment for the formation of amphiphilic structures are deep-reaching tectonic fault zones [18]. Recently, we proposed a mechanism of periodic vesicle formation which is expected to occur in fault zones filled by water and CO\textsubscript{2} [15]. At a depth of approximately -1 km, pressure and temperature conditions induce a local phase transition between supercritical CO\textsubscript{2} (scCO\textsubscript{2}) and subcritical gaseous CO\textsubscript{2} (gCO\textsubscript{2}). Various amphiphilic products of hydrothermal chemistry [19] are expected to accumulate at this point due to the solubility drop in CO\textsubscript{2} and the presence of large transition-induced interfaces.

With additional periodic pressure variations resulting from tidal influences or geyser phenomena, a cyclic process occurs in which the transition scCO\textsubscript{2} \rightarrow gCO\textsubscript{2} induces the formation of water droplets covered by a monolayer of amphiphilic compounds [20]. When migrating through the interface to the aqueous domain (which by itself is covered by a layer of amphiphiles), the droplets turn into vesicles with a bilayer membrane [21]. Being thermodynamically unstable, the vesicles are expected to disintegrate and release their organic contents into the bulk water phase over time. During the transition gCO\textsubscript{2} \rightarrow scCO\textsubscript{2}, the organic constituents and the water again become soluble in the CO\textsubscript{2} phase and the cycle can start again [15]. So, in general, each pressure cycle corresponds to one generation of vesicles, even though individual vesicles may survive for several cycles.

In the same hydrothermal environment, amino acids are expected to occur [22-24]. Under the given temperature and pressure conditions and in presence of the water/carbon dioxide interface, these amino acids undergo spontaneous condensation reactions and form a series of oligopeptides [16,25]. After a short period of time, the competing processes of condensation and hydrolysis will lead to an equilibrium situation. In this state, the concentrations of longer oligopeptides are very small. In a corresponding laboratory experiment, they decrease by approximately one order of magnitude for each additional amino acid unit [16]. Nevertheless, this process leads to a constant presence of random oligopeptides of variable length.

However, if a given peptide with a specific amino acid sequence is capable of interacting with the bilayer membranes of the vesicles described above, e.g. by being amphiphilic with an amphiphilicity profile resembling the one of the membrane, it will integrate into the bilayer structure. Such an integrated peptide is now protected against hydrolysis and therefore will accumulate over time. Consequently, its concentration can keep on growing and may surpass the original equilibrium concentration by several orders of magnitude [16]. At the same time, primarily hydrophilic peptides are recycled and primarily hydrophobic ones will be eluted by scCO\textsubscript{2} (Figure 1).
Figure 1. Mechanism of peptide selection and accumulation in presence of vesicles. Left: Peptide chains formed by hydrophilic amino acids (blue circles) will undergo little interaction with vesicles and remain in the aqueous phase where they undergo hydrolysis. Right: Peptide chains formed by hydrophobic amino acids (red circles) will eventually be eluted by scCO$_2$. Center: Amphiphilic peptides will accumulate in the bilayer membrane and remain partially protected against hydrolysis and elution [16].

Of course, such an accumulation has consequences for the vesicle structure. If the vesicle is being stabilized by the given peptide, its lifetime will increase, maybe even over several pressure cycles. This given, the period of protection for the corresponding peptide will also increase, giving this peptide further selection advantage over competing peptides. This mutual effect (peptide stabilizes the vesicle – the vesicle stabilizes the peptide) has the capability to drive an ongoing evolution of a peptide-vesicle system, targeting vesicles with an optimized potential to survive the given pressure-cycling conditions. The resulting vesicle system could be the starting point for the subsequent development of a living cell [26-28].

In the following, we want to report on an experiment which is meant to promote such an evolution process. It combines the cyclic formation and destruction of vesicles with the conditions of random peptide formation. It involves selection pressure on the vesicle structure and the observation of the optimization process over time. Finally, a resulting peptide-vesicle system is being studied for its characteristic features regarding vesicle stability and its physical properties. Overall, this experiment may be a rare example for a system which develops from a simple towards a much more complex one.

2. Materials and Methods

2.1 Amphiphiles

The choice of amphiphiles was driven by two motivations: i) focus on simple chemical structures which could easily develop in a hydrothermal system, ii) allow for vesicles which are stable at high temperatures, at a broad pH range and in presence of bivalent cations. For the two latter conditions, T. Namari and D. Deamer have proposed a mixture of long chain amines with long chain fatty acids [29] which are accessible by Fischer-Tropsch chemistry [4]. In order to improve temperature stability, C$_{18}$ chains have been chosen for both components. Accordingly, octadecylamine (GC purity ≥ 99.0%) and octadecanoic acid (GC purity ≥ 94.5%) were purchased from Sigma Aldrich.
2.2 Amino acids

The choice of amino acids again followed the condition of being accessible by hydrothermal chemistry. Initially, only proteinogenic amino acids in their natural L-form were considered for simplicity. With that, the selection was limited to a set of 12 L-amino acids which were experimentally accessible under simulated hydrothermal conditions [22]. This includes the hydrophilic amino acids glycine, serine, threonine, aspartic acid, glutamic acid, and lysine, as well as the less hydrophilic or hydrophobic varieties alanine, proline, valine, leucine, isoleucine, and phenylalanine. All amino acids (HPLC or titration purity ≥ 98.0%) were purchased from Sigma Aldrich and were used without further purification.

2.3 Pressure cell and initial setup

In order to simulate conditions given in depths between 1 and 7 km, the high pressure cell (50 mL) of a custom-made phase equilibrium apparatus (SITEC-Sieber Engineering AG, Ebmatingen, Switzerland) is used as a reaction container. It allows for pressures of up to 1000 bar with manual fine adjustment, elevated temperatures and constant stirring. It is filled with 25 mL of water (Millipore Milli-Q water, resistivity 18.2 mΩ·cm) and 25 mL of carbon dioxide (Air Liquide, GC purity ≥ 99.995%), the latter being either in the gaseous or in the supercritical state depending on the pressure and temperature conditions. A system of valves and pressure regulations allows one to add or to remove samples from both phases without changing the pressure inside the cell. All amphiphiles and amino acids were added to the aqueous phase prior to the start of the experiment. The concentrations of the amphiphiles octadecylamine and octadecanoic acid were adjusted to 0.01 M each, all 12 amino acids were added in 0.067 M concentration in the aqueous solution.

2.4 Evolution experiment

In order to accelerate the peptide formation cycle and to induce selection pressure on the vesicles, the temperature inside the cell is kept at 120°C during the whole experiment. The pressure is repeatedly switched between 100 bar and 70 bar on a regular time scale (every 30 min). During each pressure cycle, a phase transition from supercritical scCO₂ to gaseous gCO₂ and vice versa is induced. In each cycle, the transition towards gCO₂ is accompanied by the appearance of micrometer-sized droplets which, in contact with the bulk aqueous phase, form vesicles (see [15] for micrographs). On the other hand, the transition towards scCO₂ leads to a depletion of amphiphiles in the aqueous phase and therefore to (at least partial) vesicle disintegration. Cyclic pressure changes are induced by a counterbalance piston directly connected to the cell. The total evolution experiment is run for 160 h and involves 85 pressure cycles. Samples of the aqueous phase (300 µL) are taken at t=0, t=16 h, t=44 h, t=90 h, and t=160 h. Due to the presence of CO₂ at high pressure, the aqueous solution is quite acidic with a pH around 3.

2.5 PFG-NMR experiments

The PFG-NMR experiments basically follow a scheme which has been applied in earlier studies on vesicle dispersions [30, 31]. All corresponding ¹H diffusion experiments are run on a 500 MHz...
DRX spectrometer (Bruker BioSpin GmbH, Rheinstetten, Germany) with a Bruker DIFF30 probe head (1200 G/cm maximum field gradient). All measurements were performed at 298 K in a 5 mm Shigemi NMR sample tube adding 10% D2O for spin locking. As a pulse program, the stimulated echo pulse sequence is combined with two gradient pulses. A total number of 16 scans is used for each measurement. The spacing ∆ between the two gradient pulses is set to 25, 50 and 100 ms. The gradients are adjusted to strengths G between 1.5 G/cm and 750 G/cm, the gradient pulse duration δ is set to 2.0 ms. All resulting echo intensities are plotted logarithmically against the parameter γ²G²δ²(∆-δ/3) with γ being the gyromagnetic ratio of the hydrogen (¹H) nucleus. In these so-called Stejskal-Tanner-plots, each slope corresponds to the negative diffusion coefficient of the observed system component.

In order to observe the development of the vesicle properties over time, the experiments are repeated over extended time periods. In order to study the effect of the peptide, all measurements are compared with results on original vesicles. The thermal stability of the vesicles with and without peptides is assessed by introducing storage intervals at elevated temperature between the measurements (T= 50°C).

2.6 Identification of accumulated peptides

Each sample (300 µL, see section 2.4) was centrifuged in order to separate dispersed solid products from the aqueous solution, basically consisting of vesicles. The separated solid products were quantified and dissolved in 150 µL isopropanol. Prior to analysis, these samples were diluted 1:1 (v/v) with HPLC-water and analyzed along with blank samples (isopropanol/water; 1:1; v/v) for background subtraction. Both the sample and the blank were analyzed three times. Global peptide analysis was performed on an Agilent 1290 Infinity liquid-chromatography system consisting of a 1290 Infinity binary pump (G4220A) with a Jet Weaver V35 mixer, a 1290 Infinity HiP sampler (G4226A), a 1290 Infinity Thermostated Column compartment (G1316C) coupled to an IM-qTOF-MS (Agilent 6560). Separation was performed on a C18 reversed phase column (Aeris Peptide XB-C18; 150 x 2.1 mm; 1.7 µm; Phenomenex Aschaffenburg, Germany) using water and 0.1% formic acid as eluent A and acetonitrile acidified with 0.1% formic acid as eluent B. The gradient started at 5% B for 2 min, linear to 85% B after 20 min, linear to 95% B after 25 min and hold 95% B for 10 min. The column was re-equilibrated at the initial conditions for 10 min. The total run time was 45 min at a flow rate of 0.1 mL/min, the injection volume amounted to 10 µL. The column was kept at room temperature. LC flow was introduced into the IM-qTOF-MS using a dual Agilent Jet Stream Electrospray Ionization (AJS ESI) source operating in the positive mode. Source parameters were as follows: capillary 5000 V, nozzle voltage 500 V, nebulizer gas 20 psi, sheath gas 12 L/min (both N2), dry gas temperature was set to 200 °C and sheath gas temperature to 325 °C. Spectra were recorded in q-TOF only mode between m/z 50-3200.

For molecular feature finding, the MassHunter Qualitative Analysis (B.07.00) was used. Background ions from blank analysis were removed from the resulting feature list using Mass Profiler Professional (12.6.1) to avoid false positive results. Subsequently, feature list was searched for possible peptides using an in-house software tool.
MS\textsuperscript{n} analysis of possible peptides was carried out on a Merck-Hitachi D-7000 HPLC system equipped with an L-7100 quaternary pump and an L-7200 autosampler coupled to a Bruker Amazon Speed iontrap MS (Bruker Daltonics, Bremen, Germany). LC conditions were the same as described above except the flow rate of the mobile phase, which was changed to 0.15 mL/min. ESI parameters were as follows: capillary 5000 V, end plate offset 500 V, nebulizer gas 20 psi, dry gas 3 L/min (both N\textsubscript{2}) and dry gas temperature was set to 200 °C. The MS\textsuperscript{n} experiment was carried out by selecting the precursor ion of m/z 864.5 and scanning the resulting fragment ions from m/z 200 to 875 after activation with an amplitude of 1.0.

2.7 Peptide synthesis and vesicle reconstruction

The peptide NH\textsubscript{2}-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH (being the largest of the accumulated species) was synthesized commercially by APeptide Co., Ltd (Shanghai, China). The total amount of 7 mg was used in portions to reassemble the vesicle structure which has led to its accumulation. In each experiment, 2.6 mg of the peptide were added to 100 µL of a vesicle dispersion formed by a 1:1 mixture of 20 millimolar solutions of octadecanoic acid and octadecylamine at pH=3 (adjusted with 1N HCl) after 6 cyclic pressure cycles in the high pressure cell. Subsequently, the resulting vesicle dispersion was submitted to field gradient NMR experiments in order to study their average size, membrane properties and thermal stability.

3 Results

3.1 Vesicle formation

Experimental evidence for the vesicle formation during the pressure cycling is gained from optical microscopy and PFG-NMR [15,16]. The volume-averaged diameter of the vesicles amounts to 600 nm under the given circumstances, but generally depends on the rate of decompression. Some of the vesicles obviously show multilamellar structure and contain internal vesicles, both features possibly deriving from the fusion of droplets in the gas phase [15,16]. The vesicle structure formed by bilayers of amphiphilic molecules is further supported by PFG-NMR measurements which show the presence of encapsulated water molecules inside membranes of relatively low permeability (Figure 2).
Figure 2. Stejskal-Tanner-plot of PFG-NMR data obtained on water molecules in a sample of the aqueous phase during an evolution experiment. The steep initial part of the echo decay corresponds to free water, the following shallow part to encapsulated water inside the vesicles. The values for $\Delta$ (25, 50 and 100 ms) refer to the spacing between the gradient pulses.

In this set of plots, the initial, very steep decay marks the free water molecules, the negative slope corresponds to the self-diffusion coefficient of bulk water. The shallow part of each plot refers to those water molecules which are encapsulated inside the vesicles. The negative slope of this part is in accordance with a self-diffusion coefficient deriving from the Brownian motion of the vesicles. On average, it amounts to $7.1 \cdot 10^{-13}$ m²/s which, assuming the viscosity of water, corresponds to a vesicle diameter of approximately 600 nm. The PFG-NMR data obtained on the samples from the aqueous phase document the presence of vesicles over the full duration of the evolution experiment. Within its given period of 160 h, the diffusion constant as well as the vesicle diameter show a random variability of ±10%.

3.2 Peptide formation

During the same course of regular sampling, the peptides are followed by two-dimensional liquid chromatography and mass spectrometry. A principal component analysis (Figure 3) on samples taken after 0 h and 160 h reveals the presence of a fraction of molecules which i) is not present at the beginning of the experiment ($t = 0$ h), ii) is not present in absence of vesicles, and iii) is not present in raw materials (solvents, eluents). Consequently, this fraction of molecules (red frame in Fig. 3) only forms in presence of vesicles and over time and therefore should contain all species which accumulated during the evolution process.
Figure 3. Principal component analysis on samples from an evolution experiment taken after 0 and 160 h. The red box labels the entity of molecules which are absent at t = 0 h and only form in presence of vesicles over time.

All samples taken during the experiment are carefully analyzed for oligopeptides formed by the 12 amino acids listed in section 2.2. The analyses not only reveal the amino acid composition, but also allow for a rough estimation of the peptide concentrations. Selecting the species with the steepest concentration increase over time, the following oligopeptides, all belonging to the fraction “160 h with vesicles” (Figure 3), have been identified:

- Thr Thr Pro
- Lys Pro Pro Phe
- Lys Lys Gly Pro Ala
- Lys Ser Pro Ala Phe
- Lys Pro Gly Gly Gly Phe
- Lys Ser Pro Pro Ala Ala Phe Phe

The compositions given above do not represent the actual sequence of the amino acids. Instead, the amino acids are listed according to their polarity (most polar ones first). As a striking feature, the amino acid lysine occurs in almost all of the peptides. This may be related to its possible function as a charged head group: offering a free amino residue, it is expected to be almost quantitatively protonated in the given acidic environment (see section 2.4) and therefore will carry at least one positive charge. Another interesting property of the longer peptides is the abundance of hydrophobic amino acids. With 60-80% occurrence, they form the largest section of the peptide chains. Most peptides also contain phenylalanine, which is the most hydrophobic amino acid within the given selection. Altogether, one can postulate that all given peptides at least have the potential to be strong amphiphiles.
Regarding its potential to integrate into vesicle membranes and to take influence on the vesicle properties, the last species (Lys Ser Pro Pro Ala Ala Phe Phe) appears to be the most promising candidate. Therefore, this octapeptide was submitted to a closer analysis of its amino acid sequence using MS\(^{+}\) analysis with an iontrap-MS. Due to the low concentration of the peptide, the identification of the sequence was complicated and included some plausible considerations. According to the resulting data, the charged lysine residue occurs at the amino-terminated end of the peptide chain, most likely followed by the serine. At the carboxylic end of the chain, we most probably deal with a set of two alanine segments. That leaves two phenylalanine and two proline residues for the inner part of the chain. Due to the relatively bulky phenyl side group, it is less likely that two phenylalanine units are directly connected. Proline, on the other hand, induces a relatively stiff chain conformation and has the property to induce turns in the peptide chain. Therefore, two prolines are unlikely to be directly connected as well. That leaves an inner sequence of either Pro Phe Pro Phe or Phe Pro Phe Pro for the central part of the octapeptide. Based on these findings and conclusions, we decided for

\[
\text{NH}_2\text{-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH}
\]

as the most likely structure of the given peptide. In the individual chromatogram of the peptide (Figure 4), three peaks can be observed which represent the same composition of the peptide, but obviously different sequences. All three peptides have accumulated in the evolution experiment. Therefore, we can conclude that there is a certain variability in the sequence and that the sequence given above is most likely among those accumulated species.

![Figure 4: Extracted ion chromatogram of the selected mass peak representing the octapeptide composition](image)

3.3 Vesicle reassembly

Having identified an accumulated species, the main purpose of this study is to elucidate why this particular octapeptide had the potential to be selected during the evolution experiment. This
question is efficiently approached by a study on the special properties of a corresponding peptide-vesicle system. Since the concentration of the octapeptide in the overall mixture is still extremely low, this task has to be performed on a reconstituted system. For this purpose, the octapeptide NH₂-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH is commercially synthesized and added to a neat vesicle dispersion which has been prepared in the high pressure cell. The resulting peptide-vesicle system is carefully studied and compared to the original vesicle system.

When added to the original vesicles, the peptide has an immediate influence on the vesicle size (Figure 5). The volume-averaged hydrodynamic radius of the vesicles is determined from the slope of the shallow part of the Stejskal-Tanner plots. This slope corresponds to the negative diffusion coefficient associated to the Brownian motion, which in turn allows for the calculation of the vesicle size at a given temperature and solvent viscosity. From an average value of 600 nm, the diameter is suddenly reduced to 300-400 nm by the action of the peptide. Before and after the addition of the peptide, the diameter is relatively stable over time.

![Figure 5: Average diameter of the vesicles before (left) and after the addition of the peptide NH₂-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH (right of dotted line). The error bars mark the experimental variability.](image)

A special feature of the PFG-NMR measurement and of the resulting Steijskal-Tanner plot is its sensitivity for the permeation process. The variation of the level of the shallow part with the gradient pulse spacing Δ allows for direct conclusions on the membrane permeability for the observed molecules (water in the given case). Due to the exchange-related loss of encapsulated water molecules over time, the permeation process leads to a corresponding drop of the final level of the plot with increasing pulse spacing Δ. In case of the neat vesicles, this drop is not observable. Instead, there is even an inverted sequence (the level for 100 ms is higher than the one for 25 ms), which results from internal water diffusion inside of the vesicle volume (Figure 6, top row). However, as soon as the peptide is added, the sequence changes drastically. Actually, the sequence now is typical for a rapid permeation process: the level for 25 ms is higher than the one for 100 ms (Figure 6, first image in the bottom row). Obviously, there is a significant loss of encapsulated water molecules in the time interval between 25 and 100 ms which is induced by the addition of the peptide.
Figure 6: Steijskal-Tanner plots determined on water for neat vesicles (top row) and after the addition of the peptide NH₂-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH (bottom row). As determined from the inversion of the sequence for \( \Delta = 25, 50 \) and 100 ms, the peptide causes a temporary increase of the vesicle membrane permeability for water (bottom row, left). Within several hours, the sequence slowly returns to the original one, indicating the loss of the original permeability over time (bottom row, from left to right).

However, this situation changes over time. Within 1.5 h, the permeation-induced drop collapses, and gradually returns to the original one observed on neat vesicles (Figure 6, bottom row, left to right). Finally, after 108 h storage at elevated temperature (50°C), the original sequence is reached (Figure 7, bottom row, right).

The most interesting aspect of the peptide selection is the potential influence on the thermal stability of the vesicles. For the assessment of the given peptide, the PFG-NMR measurements have been repeated during a period of high-temperature storage (50°C, 100h), for neat vesicles as well as for the peptide-vesicle system. The result is shown in Figure 7. It reveals a significant thermal instability of the neat vesicles, as determined from the drop of the final plateau values (Figure 7, top row). The loss of the encapsulated fraction corresponds to a half-life time around 100 h. In contrast, no such loss is observed for vesicle dispersions after the addition of the peptide (Figure 7, bottom row). This demonstrates a strong stabilizing effect on the vesicles which is induced by the selected octapeptide NH₂-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH.
4 Discussion

The experimental results clearly show a significant fraction of peptides which i) are not yet present at the beginning of the experiment and gradually form over time, and ii) only form in presence of the vesicles. Among these, a few peptides are sticking out in terms of their concentration increase over time, most prominently the octapeptide NH₂-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH. Obviously, this peptide has been accumulated in a selection process induced by the presence of membrane vesicles. Looking at possible mechanisms of peptide selection in detail, one can differentiate between three possible criteria favoring different types of amphiphilic peptides (Figure 8) [16]. In the following, we briefly describe the three selection criteria and discuss the possible contribution of the selected peptide NH₂-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH:

Figure 7: Stejskal-Tanner plots determined on water for neat vesicles (top row) and after the addition of the peptide NH₂-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH (bottom row). Both series correspond to storage at 50°C for 0, 19 and 100 h. As determined from the shift of the final plot level, there is a significant decomposition of the neat vesicles (top row). No such decomposition is observed on vesicles containing the peptide, revealing its stabilizing effect.
Figure 8. Possible targets of a selection process of peptide-vesicle systems. Integration and protection of peptide (parasitic), additional thermal stabilization of the vesicle structure (symbiotic), and introduction of a peptide-induced function (functional).

1) Integration. Amphiphilic peptides with an amphiphilicity profile reflecting the one of the bilayer integrate into the vesicle membrane. Hereby, they gain an individual selection advantage as they are partially protected against hydrolysis and as they become less easily eluted from the vesicle zone. For this selection criterion, the effect of the peptide on the stability of the vesicle is irrelevant, hence one could call this mechanism a parasitic one.

The amphiphilicity profile of the selected peptide NH₂-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH is clearly predestinated by its sequence. In the given acidic environment (pH=3), the initial lysine group, offering an additional amino residue, will carry at least one positive charge, possibly even two since it forms the amino end of the chain. Serine is quite hydrophilic as well and can therefore contribute to the polar head of the molecule. The residual chain of the peptide is being formed by six non-polar amino acids (pairs of proline, phenyalanine, and alanine) with a clear potential to represent the hydrophobic part of the molecule. Altogether, the selected octapeptide can be regarded as strongly amphiphilic. All experimental results support its expected rapid membrane integration. The changes of the vesicle size (Figure 5) and vesicle membrane permeability (Figure 6) occur almost instantaneously and can only be interpreted by an intense and integrative peptide-membrane interaction.

2) Stabilization. In this “symbiotic” interaction, amphiphilic peptides are again being protected by vesicles, but in turn they also stabilize the vesicle structure. This leads to a mutual advantage connected to the formation of the peptide-vesicle system. With increased stability, the vesicles could even survive several pressure cycles (“generations”), therefore giving the peptide an increased degree of protection over a longer period of time. Consequently, this “symbiotic” effect would lead to an even stronger selection advantage of the peptide.

In the given experiment, the selected peptide NH₂-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH clearly has a protective effect on the vesicles. While the original vesicles show a significant thermal decomposition over 100 h at 50°C (Figure 7, top row), the peptide-vesicle system seems to be stable under the same conditions (Figure 7, bottom row). Even though there is no plausible interpretation of the mechanism, the effect of thermal stabilization is obvious. An additional factor for the stability-related selection of vesicles may be the vesicle size. During the decompression step of the
pressure cycling, CO₂ bubbles are generated which have the potential to disrupt vesicles if they form inside their inner volume. Consequently, smaller vesicles have better chances to stay intact, giving peptides which reduce the vesicle size a selection advantage. This may explain the effect of the selected peptide on the vesicle diameter (Figure 5).

3) Function. The selection of peptides could go even beyond the effect of simple mechanical stabilization and lead to more complex (“functional”) survival mechanisms. A possible example may be a membrane structure which allows for the increased permeation of solvent through the membrane (like, e.g., by the formation of a channel structure [32]). In the vesicle cycle, the vesicles are generated with a natural ionic concentration gradient across the membrane which destabilizes their structure by the resulting osmotic stress. Permeation through the membrane could cause a rapid relaxation of this gradient, leading to a longer vesicle lifetime and resulting in a corresponding selection advantage for the permeation-inducing peptide.

Obviously, the selected peptide NH₂-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH causes an increased degree of water permeation through the vesicle membranes (Figure 6), at least for a limited period of time. This period, however, is sufficient for an equilibration of concentration gradients which mainly consist in the dissolved amino acids (their overall concentration being 0.8 M in the outer aqueous phase, but nearly zero in the inner phase of the vesicles). Again, there is no clear interpretation of the mechanism and the reason why it only acts temporarily, but the effect of water permeation is significant and will probably include dissolved components as well. It will lead to a rapid decrease of the osmotic pressure and therefore increase the survival rate of the vesicles.

5 Conclusion

The peptide NH₂-Lys-Ser-Pro-Phe-Pro-Phe-Ala-Ala-OH selected in the described experiment may be an example for the result of a molecular evolution process occurring under very primitive and quite natural conditions. With the given sequence, it obviously integrates into the vesicle membrane, increases its thermal stability as well as its permeability and decreases the vesicle size. All these effects can be interpreted as suitable survival strategies of the peptide-vesicle system leading to prolonged lifetimes under the given circumstances.

Even though the described mechanism of peptide selection misses the capability for identical reproduction, the mechanism of selection from a large pool of random peptides over a long period of time can be quite efficient. Both conditions are definitely fulfilled for peptides in hydrothermal sources. The entropic driving force of the process consists in the expansion and dilution of a large amount of hydrothermal products rising in the Earth’s crust, from which a small fraction is selected over a long and iterative process (Figure 9). The outcome may be a peptide-vesicle system which has developed a set of functions leading to its long-term survival. In the evolution process on early Earth, this may have included an early stage of a primitive metabolism which could have taken advantage of concentration gradients as an initial energy source by the use of catalytically active peptide channels. Finally, such a functional peptide-vesicle system could also have formed an ideal platform for a subsequent RNA world.
Figure 9. Schematic diagram of a possible structural evolution of vesicles by repeated optimization steps in tectonic fault zones.

References

Black, R.A., Blosser, M.C., Stottrup, B.L., Tavakley, R., Deamer, D.W., Keller, S.L. Nucleobases bind to and stabilize aggregates of a prebiotic amphiphile, providing a viable mechanism for the emergence of protocells. PNAS 2013, 110, 13272-13276.


Mayer, C., Bauer, A. Molecular exchange through capsule membranes observed by pulsed field gradient NMR. Prog Coll Polym Sci 2006, 133, 22-29.
