INTRODUCTION

S100 proteins are Ca\(^{2+}\)- and Zn\(^{2+}\)-binding proteins that constitute the largest subgroup within the superfamily of Ca\(^{2+}\)-binding EF-hand proteins. In humans, 21 members have been identified so far [1,2]; these show a tissue- and cell-type-specific expression pattern and characteristic subcellular localizations [3]. S100 proteins became the focus of intensive research owing to their association with numerous human disorders, including acute and chronic inflammatory conditions, autoimmune diseases, cancer, atherosclerosis, cardiomyopathies and neurodegenerative diseases [4-8], apart from their crucial roles in cell physiology. These proteins are able to perform a plethora of intra- and extracellular functions, including cytokine-like and chemokine-like activities via activation of the receptor for advanced glycation endproducts (RAGE) [9] and toll-like receptor 4 (TLR4)-dependent signalling cascades [10], regulation of the cytoskeleton via tubulin polymerization [11] and other functions. Recently, a new property among S100 proteins was unveiled: we have discovered that the S100A8/A9 proteins form amyloids both in the aging prostate and in vitro [12]. Amyloidogenic propensity in vitro is also shared by other S100 proteins [13], including S100A6 [14]. Here, we review and discuss the conformational plasticity of S100 proteins, which may underlie their multiple pathways of self-assembly leading to their multiple roles in health and disease.

Some abbreviations are listed at the end of the chapter.
STRUCTURAL DIVERSITY OF S100 MONOMERS AND MULTIMERS

S100 Structure and Folding

S100 proteins are rather small proteins with a length of 79 to 114 amino acid residues corresponding to a size of 9 to 14 kDa [1,15,16]. Their sequence identities range from 22% to 57%, showing that the diversity within the family is quite large (Fig. 18.1) [1]. Under physiologic most S100 proteins form multimers ranging from dimers to decamers (Fig. 18.2) [17–19]. Low salt concentrations stabilize monomers which retain the native fold [20]. The basic structure of the different S100 protein multimers is very similar. Each monomer comprises two adjacent EF-hands connected by a flexible linker (Fig. 18.2A). The N-terminal EF-hand, composed of helices I and II, is specific for S100 proteins whereas the C-terminal EF-hand, involving helices III and IV, shows the classic architecture present in all other EF-hand proteins. Dimerization of S100 proteins occurs via helices I and IV from both monomers, forming a stable four-helix bundle. Helices II

![FIGURE 18.2] Multiple sequence alignment of human S100 proteins. The sequence identities among different S100 proteins range from 22% to 57%. The highest homology is observed in the EF-hand regions. Ca²⁺ coordinating residues in the EF-hand motifs are highlighted in red. Bright red indicates residues which coordinate Ca²⁺ via their backbone oxygen; dark red highlights residues which coordinate Ca²⁺ via side-chain oxygen. Zn²⁺ coordinating residues identified in structural studies are highlighted in blue. The largest variations in sequence are observed in the so-called hinge region, between helices III and IV, and in the C-terminus.
and III of both monomers reside on each side of the dimer. These dimers serve as building blocks for the formation of larger multimeric assemblies such as tetramers, hexamers, octamers or even decamers (Fig. 18.2B–F). Most of these assemblies are homo-multimers. However, several S100 proteins form hetero-multimers as well, such as e.g. S100A8 with S100A9, S100B with S100A1 or S100A6 or S100A11 [21]. Multimerization is very often triggered by Ca\(^{2+}\) or Zn\(^{2+}\) binding, as in the case of S100A8/A9 heterotetramer formation (Fig. 18.2C) [19,22] or of S100A12 hexamerization (Fig. 18.2D), where both metal ions play a crucial role [23].

Ca\(^{2+}\) and Other Metal-Binding Properties

Many S100 proteins act as Ca\(^{2+}\) sensors, translating changing intracellular Ca\(^{2+}\) concentrations into a signal. In the inactive Ca\(^{2+}\)-free state, the N-terminal EF-hand harbors a Na\(^{+}\) ion that is replaced by Ca\(^{2+}\) when Ca\(^{2+}\) levels rise beyond a certain threshold [24,25]. In this EF-hand, Ca\(^{2+}\) is coordinated mainly by backbone carbonyls and exhibits lower Ca\(^{2+}\) affinity than the classic C-terminal EF-hand, where Ca\(^{2+}\) is mainly coordinated by side chains from aspartate, asparagine and glutamate residues [16]. Mutation of one of these side chains results in the loss of Ca\(^{2+}\) binding as observed e.g. in S100A10. Ca\(^{2+}\) binding to S100 proteins triggers a large conformational change, opening the structure by a 90° degree movement of helix III [26]. The hydrophobic target interaction site that is exposed by the conformational change is mainly formed by residues from helix IV and the linker region [15].

Many S100 proteins bind Zn\(^{2+}\) with high affinity at sites distinct from the Ca\(^{2+}\) ones. The Zn\(^{2+}\)-binding S100 proteins can be divided into two groups. In the first group,
comprising S100A2, S100A3 and S100A6, coordination of Zn$^{2+}$ involves predominantly cysteine residues [27–30]. In the second group, histidine, glutamate or aspartate residues bind Zn$^{2+}$ ions; this group includes S100A7, S100A7A, S100A8, S100A9, S100A12 and S100B. Interestingly, Zn$^{2+}$ binding to S100A12 or S100B increases the affinity of the proteins towards Ca$^{2+}$ by up to 1500-fold [31-33]. Moreover, in the presence of Zn$^{2+}$ the affinities of S100A9 and S100A12 towards their receptors RAGE or TLR4 increase by several hundred-fold [23,34], revealing a further regulatory function of Zn$^{2+}$.

Cu$^{2+}$ binding has been reported for S100B, S100A12 and S100A13 [35–38]. However, the in vivo free Cu$^{2+}$ concentrations are rather low, raising the question of the physiologic role of the Cu$^{2+}$ binding observed in vitro. Recently, high-affinity binding of Mn$^{2+}$ has been observed for S100A8/A9 and revealed a new function of this protein [39]. As a part of the innate immune response, S100A8/A9 is released in high concentration at places of pathogenic invasion where it limits the availability of trace metal ions essential for microbial growth. Binding of Mn$^{2+}$ effectively inhibits the growth of pathogens and weakens their defense towards reactive oxygen species [40].

### TABLE 18.1 Influence of Metal Ions on the Oligomeric State and Function of Some S100 Proteins

<table>
<thead>
<tr>
<th>S100 Protein</th>
<th>Oligomerization State</th>
<th>Is Oligomerization Metal-Dependent?</th>
<th>Function</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>S100B</td>
<td>Dimer</td>
<td>No</td>
<td>Binds RAGE</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>Tetramer</td>
<td>No</td>
<td>Binds RAGE with higher affinity</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hexamer/octamer</td>
<td>Ca$^{2+}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>S100A12</td>
<td>Dimer</td>
<td>No</td>
<td>Binds RAGE</td>
<td>43</td>
</tr>
<tr>
<td></td>
<td>Hexamer</td>
<td>Ca$^{2+}$</td>
<td>Binds RAGE with higher affinity</td>
<td></td>
</tr>
<tr>
<td>S100A4</td>
<td>Multimeric</td>
<td>No</td>
<td>Neurite outgrowth</td>
<td>45</td>
</tr>
<tr>
<td>S100A8/A9</td>
<td>Heterodimer</td>
<td>No</td>
<td>Microtubule formation</td>
<td>46</td>
</tr>
<tr>
<td></td>
<td>Heterotetramer</td>
<td>Ca$^{2+}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>S100A2</td>
<td>Dimer</td>
<td>No</td>
<td>Tumor suppressor</td>
<td>47</td>
</tr>
<tr>
<td></td>
<td>Tetramer</td>
<td>Zn$^{2+}$</td>
<td></td>
<td>29</td>
</tr>
</tbody>
</table>

Apart from Ca$^{2+}$, Zn$^{2+}$ was also reported to influence the degree of oligomerization of S100A2. The binding of Zn$^{2+}$ to its lowest affinity binding site induces the formation of an S100A2 tetramer via the assembly of two homodimers, whereby the metal is coordinated by a cysteine from S100 OLIGOMERS, AGGREGATES AND AMYLOIDS

### S100 Functional Oligomers

Multimer formation most likely regulates the activity and modifies the function of S100 proteins (Fig. 18.2). For multimerization – as well as for conformational changes and stabilization – metal ions such as Ca$^{2+}$ and Zn$^{2+}$ have an important role (Table 18.1) [13]. For example, S100B forms larger oligomeric species such as tetramers, hexamers and octamers (Fig. 18.2E). The formation of the two largest multimers is enhanced by Ca$^{2+}$ binding, while the addition of EDTA eliminates these assemblies [18]. The multimers represent a large portion of S100B in human brain but the most stable is the tetramer, which does not dissociate into dimers [18]. Binding of dimeric and tetrameric S100B to RAGE is strictly Ca$^{2+}$ dependent. However, tetramers bind RAGE with higher affinity and promote cell survival whereas dimeric S100B binds with lower affinity and triggers a pro-apoptotic response [18].

S100A12 also requires Ca$^{2+}$ for interacting with its targets, including RAGE. The RAGE–S100A12 interaction mediates a pro-inflammatory response to cellular stress and can contribute to the development of inflammatory lesions [41,42]. As for S100B, the S100A12 hexamer is formed in a Ca$^{2+}$-dependent manner (Fig. 18.2D) and has enhanced binding affinity to RAGE [43]. Another recent example is S100A4, where a specific inhibitor induces the formation of a homodecamer (Fig. 18.2F) which cannot bind target proteins any more, indicating that multimer formation is a potent modulator of S100 protein activity [44]. In addition, extracellular S100A4 oligomers are potent promoters of neurite outgrowth and survival [45]. Finally, the Ca$^{2+}$-induced S100A8/A9 heterotetramer (Fig. 18.2C) controls microtubule formation [46].
Each subunit [29]. Metal ions thus play an important role in the formation of functional S100 oligomers, and recent observations on S100A8/A9 [12,13] suggest that they may also be involved in the formation of oligomeric and fibrillar assemblies.

Amyloidogenic and Aggregation Potential of S100 Proteins

Some members of the S100 protein family possess an inherent potential to form amyloid fibrils. Despite the fact that several factors influence the amyloidogenic process, the primary sequence — to a large extent — defines the aggregation and fibrillation propensity of a polypeptide chain. This potential can be estimated through different computational algorithms, e.g. Zyggregator, that estimates the aggregation propensity [48], and Waltz, which is able to identify amyloid-forming regions in functional amyloids [49]. Indeed, the intrinsic aggregation propensity scores of the monomeric forms of S100A8 and S100A9 at pH 2.5 and 7.0 were determined to be of the same order of magnitude as that of the Aβ peptide, which forms amyloid deposits in Alzheimer’s disease [12]. Taking into account that S100 proteins have a similar chemical and structural identity, it was hypothesized that the ability to form amyloid structures could be a generalized property among the S100 protein family. When other S100 proteins, including S100A3, S100A6, S100A12 and S100B, were tested for the formation of amyloid-like structures at pH 2.5 and 57°C, it was found that most of them formed thioflavin-T (ThT)-binding amyloid structures, with S100A12 being the only exception. The formation of both amyloid fibrils and other precursor structures was confirmed by atomic force microscopy (AFM) [13].

Effects of S100 on Aggregation Cross-Seeding

Although several S100 proteins aggregate into amyloid-like structures, some of them have the ability to modify or enhance the aggregation potential of other proteins. An example is S100A6, an amyloidogenic protein that aggregates into fibrils under physiologic conditions and is capable of enhancing the aggregation of SOD1 [14]. Both proteins are involved in amyotrophic lateral sclerosis (ALS), a fatal neurodegenerative disorder [50]. The Zyggregator algorithm predicts a high propensity for the formation of amyloid-like structures by S100A6 [14]. In fact, at acidic conditions and high temperature (pH 2.5 and 57°C) the ThT binding kinetics revealed a fast fibril formation process with no lag phase. The formation of aggregates and fibrils was also observed through transmission electron microscopy (TEM). Interestingly, Ca²⁺ produced an inhibitory effect on the aggregation kinetics, probably because this metal ion promotes anti-parallel β-sheet conformations that can repress fibrillation. Under physiologic conditions (pH 7.0 and 37°C) this protein also forms amyloid-like fibrils in a slower process, but without a lag phase (Fig. 18.3A). The formation of S100A6 amyloidogenic structures lead us to hypothesize a new role for S100 proteins in brain damage. S100A6 is overexpressed in astrocytes which are associated
with degenerating motor neurons in ALS patients. In these neurons, SOD1-enriched inclusions were also found [51], rendering the amyloid cross-seeding between the two proteins a real possibility under in vivo conditions [52]. Botelho et al showed that native S100A6 nucleates SOD1 fibrillation, leading to a decrease in the lag phase by more than 10 h (Fig. 18.3B), thus suggesting a novel role for S100A6 aggregation in human neuropathologies [14].

The pro-inflammatory protein S100A9 was also shown to interact with the Aβ1-40 peptide, inducing its fibrillation [53]. Co-incubation of S100A9 and Aβ1-40 at physiologic conditions (pH 7.4 and 37°C) produced massive fibrillar aggregates observed by AFM. The interaction of these two proteins was also observed through the kinetics of amyloid formation (ThT-binding assay) and transitions in secondary structure (far-UV circular dichroism). Interestingly, the addition of Aβ1-40 inhibits S100A9 cytotoxicity. This may occur via the binding of toxic S100A9 species to Aβ1-40 amyloid structures. Altogether, these results suggest that S100A9 secretion during inflammation leads to the formation of amyloid plaques. This can nevertheless be a protective mechanism against Aβ-associated toxicity in Alzheimer’s brains.

Role of Metal Ions in Aggregation Pathways

The binding of different metal ions to S100 proteins may modulate their conformation, folding and – ultimately – function. As discussed above, the identification of high amyloidogenic potential in S100 proteins suggests that they may have the ability to acquire amyloid-like conformations under certain physiologic conditions, and it becomes clear that metal ions can have an important role in this process (Fig. 18.4) [13]. The general role of metal ions as modulators of protein conformation, misfolding and aggregation in disease has been recently reviewed [52].

In the case of S100A2, binding of Ca²⁺ and Zn²⁺ promotes conformational changes that alter protein stability: Ca²⁺ has a stabilizer role whereas Zn²⁺ has the opposite effect [54]. These metal ions also have an impact on the formation of functional oligomers, as in the case of the Ca²⁺-dependent S100B hexa- and octamerization [18]. Ca²⁺ and Zn²⁺ also play a determinant role in promoting amyloid assembly of S100A8/A9 proteins in vivo [12]. Besides the above mentioned factors, metal ion dyshomeostasis or anomalous protein–metal interactions can also influence S100 functional states within a physiologic context, leading to the formation of amyloid structures or precursor oligomers [52].

**BIOIMAGING OF S100 AMYLOIDS BY AFM AND TEM**

**Techniques to Study Protein Amyloid Formation**

Protein polymerization and aggregation in vivo and in vitro is far from being understood, and numerous techniques
have been developed in order to unravel the underlying mechanism with the ultimate aim of understanding the respective pathologies and establishing inhibition strategies and therapeutic approaches. X-ray diffraction techniques, solid-state magic-angle spinning NMR spectroscopy, circular dichroism spectroscopy and microscopy studies, among others, have been utilized to detect and examine the chemical, electronic, material and structural properties of amyloid fibrils at up to Ångström spatial resolution [55]. Nevertheless, imaging amyloid morphology at the ultrastructural level is still a challenge and remains a goal of researchers.

Both AFM and TEM microscopies play a key role in studies on the morphology and pathways of amyloid assembly. Typical amyloid fibrils are characterized by a nanometer size in cross-sectional dimensions and can reach up to micrometer length and longer in their axial growth, which makes them a perfect object for these microscopy techniques [56–58], while being far too small for optical microscopy. AFM and TEM provide sub-nanometer and even up to Ångström (AFM in z-axis) resolution and at the same time can image micrometer size structures. This over 1000-fold dynamic range surpasses the optical diffraction limit attainable by optical microscopes and provides insights into sub-molecular details comparable with high-resolution molecular structural techniques. AFM does not require any special treatment of the samples and is able to image them in air or liquid. For TEM studies, biologic macromolecules, due to their low density and therefore contrast, are subjected to negative staining, rotary shadowing and cryo-EM [56].

Both TEM and AFM have been widely used for imaging a variety of materials, ranging from carbon nanotubes [59] to human tissues, and they constitute a powerful tool that even allows atomic resolution. Amyloid fibrils of different origins have also been imaged in vivo, ex vivo and in vitro by both AFM and TEM. Compared to other techniques, TEM and AFM usage is more widespread, easier to access and handle, and has been used to characterize various assembly states of Aβ, α-synuclein, IAPP, TTR and lysozyme, among others [12,57,60–65].

Sources of Amyloid – In Vivo and In Vitro

In vivo imaging of amyloids by microscopy techniques in affected tissues is limited by the available tissue sources and cannot provide information on the dynamics of fibril formation. Most studies are based on observations using extracted amyloids and fibrils produced in vitro. In vitro fibril production has been a powerful tool for studying the mechanisms leading to amyloid formation. Theories for the events occurring in vivo have been proposed based on in vitro results. Virtually all types of fibril were produced in vitro from their native precursors, suggesting that the amyloidogenic potential is a feature shared by all polypeptide chains [66]. For in vitro studies, amyloid fibrils are generated in solution or suspension and then captured on mica in air or buffer (for AFM studies) or on a rapidly prepared TEM grid. Much evidence has been accumulated showing that synthetic fibrils are morphologically similar to those formed in vivo, indicative of a predominant role for the amyloidogenic potential of proteins in amyloid disorders.

Fibril formation in vitro also opened the possibility of studying the process by which a native soluble protein loses its normal fold and aggregates into fibrils, paving the way for a better understanding of the mechanism that culminates in amyloid formation and the design of drugs that interfere with such a process. Protein denaturation and the amyloidogenic pathway involve the formation of several intermediates with variable quaternary, tertiary and secondary structures. Accordingly, several amyloidogenic intermediate species en route to mature amyloids have been identified in different amyloidotic disorders. Furthermore, fibril formation is dependent on factors such as pH, protein concentration, time and metal ions. Here we describe highly heterogeneous S100 protein aggregates visualized by microscopy techniques and occurring both in vitro (under different pH, Ca²⁺ and other salt concentrations) and extracted from ex vivo tissue.

S100A8/A9 Amyloids

The first proteins in the S100 family that were found to be amyloidogenic were S100A8/A9 [12]. Indeed, the overall aggregation scores for S100A8 calculated using intrinsic propensities for the aggregation of individual amino acids are comparable to those of Aβ [12,67]. In agreement, amyloid structures of S100A8/A9 were formed in vitro from proteins extracted from granulocytes (and produced recombinantly in Escherichia coli) which were incubated at pH 7.4, 37°C with agitation, or pH 2.0, 57°C without agitation. Under both conditions, S100A8/A9 were found to assemble into heterogeneous fibrillar species as illustrated in Figure 18.5. At pH 7.4, oligomeric species and short protofilaments were observed after 2 weeks of incubation (Fig. 18.5A), while upon further incubation, for up to 8 weeks, thick fibrillar bundles of a few micrometers in length were massively populated (Fig. 18.5B,C). Their dimensions were estimated in AFM cross-sectional analysis by measuring z-height, the most accurate AFM parameter. The height of fibrillar bundles reached ca. 15–20 nm, while the height of individual filamentous structures was ca. 5 nm (Fig. 18.5G,H). Fibrillar lateral assembly and thickening may be a contributing factor to their stability and resistance to different environmental factors and proteases, when they spontaneously assemble in vitro or in vivo.

In the S100A8/A9 samples incubated at pH 2.0 without agitation, the spherical oligomeric species and protofilaments were also formed in ca. 2 weeks (not shown).
After 4 weeks, short straight fibrils of sub-micrometer length were highly populated, as observed very consistently by both AFM and TEM imaging (Fig. 18.5 D,E). They coexisted together with long filamentous structures from a few hundred nanometers to micrometer scale lengths, which were also observed by both AFM and TEM (Fig. 18.5F). The height of representative structures in AFM cross-section was ca. 4–5 nm (Fig. 18.5F top, Fig. 18.5H).

Amyloidogenic Properties of S100A6 Studied by TEM

The association of S100 proteins – namely, their aggregated states – with human disease has been recently expanded by the description of S100A6 as an amyloidogenic protein and its seeding effect towards SOD1 amyloidogenesis, characteristic of ALS [14]. The importance of this finding was two-fold: on the one hand, it suggested...
novel mechanisms in ALS etiology; on the other hand, it pointed to widespread amyloidogenic potential in the S100 family, with as yet unknown relevance. For this reason, the high-resolution characterization of these novel amyloids and the corresponding aggregation pathways is invaluable. S100A6 forms typical amyloid fibrils under acidification (Fig. 18.6A,C,E). As reported for other amyloidogenic proteins [61–63], the fibrils are polymorphic, exhibiting different species [14]. It is noteworthy that there are fibrils of 4–5 nm diameter, isolated or laterally assembled into thicker fibrils (Fig. 18.6A, black arrows) and 15 nm wide fibrils of variable length (Fig. 18.6A, arrowheads). Other structures resembling oligomers were also visualized (Fig. 18.6A, gray arrow). Interestingly, coiled fibrils were not observed, although additional analyses, such as metal shadowing followed by TEM, are necessary to completely rule out the presence of this feature. Besides the differences in diameter, it is also interesting to note the presence of long thin fibrils and shorter thicker fibrils. This may be representative of distinct aggregation mechanisms, occurring at least in vitro. Accordingly, the long thin fibrils of 4–5 nm diameter correspond to elementary protofilaments formed by vertical stacking of the basic unit, e.g. the S100 monomer [13]. These protofilaments can

FIGURE 18.6  Effect of pH and buffer conditions in S100A6 and S100B fibril formation. (A) S100A6 fibrils assembled at glycine pH 2.5 for 28 days are polymorphic in nature, presenting 4–5 nm wide fibrils, isolated or laterally assembled in thicker fibrils (A, black arrows) and 15 nm wide fibrils of variable length (A, black arrowhead). Other structures resembling oligomers are also visualized (A, gray arrow). (C) Increasing pH to 7.0 in phosphate buffer results in single 8 nm wide fibril species distinct from the ones obtained at lower pH. (E) At pH 7.0 and Tris 50 mM, the process of fibril formation is delayed as only amorphous aggregates and pre-fibrillar material are detected. S100B also forms fibrils, but to a lower extent. (B) At pH 2.5, fibrils are generated abundantly but the morphology is closer to that of S100A6 fibrils at pH 7.0 in phosphate buffer. (D) A similar situation occurs at pH 5.0 acetate. (F) At neutral pH, S100B does not form fibrils and only intermediate species are observed, at least during 28 days incubation at 37°C.
associate with each other and form higher-order structures, constituting mature fibrils. Alternatively, the later can be formed by longitudinal growth of full-width species, justifying the presence of thicker but shorter fibrils. Indeed, this pathway might also apply to the formation of S100A6 fibrils because thick, but short, structures were observed. Whether these two distinct mechanisms exist in vivo is not known, but they may represent a modulation of S100 fibril formation, with different impacts on cellular toxicity and disease development.

At pH 7.0 (50 mM phosphate), S100A6 fibrils were still formed, although less abundantly than at pH 2.5 (Fig. 18.6C). Interestingly, increasing pH also rendered the fibrils’ population more homogenous, especially concerning the diameter, as only approximately 8 nm wide fibrils were observed. The 8 nm wide fibril is also reported as the most prominent species for other amyloid fibrils generated in vitro, such as TTR [61]. Nevertheless, it is still possible to detect fibrils of variable length and, occasionally, small oligomers (Fig. 18.6C, arrowheads). In this case, fibrils seem to form upon stacking of these spherical oligomeric species (Fig. 18.6C, arrows). The higher-order thicker fibrils are not observed under these conditions. The observation that S100A6 aggregates differently at pH 7.0 when using 50 mM tris indicates that salt concentration and ions also play an important role in the aggregation process. In this case, the fibril formation process appears to be delayed as mainly aggregates and pre-fibrillar material were detected (Fig. 18.6E).

**S100B Amyloid Formation In Vitro**

Similarly to S100A6, S100B enters different aggregation pathways at different pH values. However, the morphology of – the much more abundant – S100B fibrils generated at pH 2.5 (50 mM glycine, Fig. 18.6B) is closer to that of S100A6 fibrils at pH 7.0 (Fig. 18.6E). (In fact, increasing the pH to 5.0 and 7.0 does not result in fibril formation, as observed for S100A6, but instead leads to the formation of intermediate species, such as large amorphous aggregates and oligomers of different sizes and shapes, as those depicted in Fig. 18.6D and Fig. 18.6F, respectively). In this regard, the S100B behavior at pH 7.0 in 50 mM phosphate is closer to the behavior of S100A6 in tris buffer, indicating a slower kinetics and suggesting a lower amyloidogenic potential. The kinetics of S100A6 and S100B fibril formation is slow when compared to that of other amyloidogenic polypeptides, such as amylin and Aβ [68,69] or even other proteins with slow assembly rates, such as TTR [61]. Upon 33 days of incubation of S100A6 at 37°C, TEM showed only the formation of aggregates and oligomers (Fig. 18.7A). Even after 65 days, the presence of fibrils was scarce (Fig. 18.7B) and aggregates and oligomers were still observed abundantly.

Long incubation times, such as 84 days at 37°C, result in the production of homogenous fibril preparations (Fig. 18.7C). These fibrils were variable in length but very homogenous in diameter (8–10 nm). An exhaustive sample visualization also permitted the detection of spherical species, resembling either the monomeric protein or small oligomers about 6–7 nm in diameter. Further incubation over time resulted mainly in increased length whereas no alterations were visualized regarding fibril diameter (Fig. 18.7D). S100B shows a similar behavior, and thus the mechanism of fibril formation seems to involve the rearrangement of aggregates and oligomers (Fig. 18.7E) into distinct fibrils (Fig. 18.7F, arrow). Again, kinetics is slower than for S100A6, and after prolonged incubation at 37°C large aggregates and oligomers were also abundantly visualized, despite fibrils being present in generous amounts, (Fig. 18.7F, arrowheads).
As we have discussed above, S100 proteins are Ca\(^{2+}\) and Zn\(^{2+}\)-binding proteins, and the study of metal-modulated aggregation propensity has just started. Recent data showed that Ca\(^{2+}\), but not Zn\(^{2+}\), represses S100A6 aggregation, resulting in antiparallel β-sheet conformations, indicative of oligomeric species structurally distinct from fibrils, as assessed by Fourier transform infrared spectroscopy \[14\]. TEM analysis provided new insights on the structures formed, confirming the presence of oligomers in the presence of Ca\(^{2+}\) (Fig. 18.8B,D). Additionally, TEM also shows that the nature of the oligomers formed in the presence of the ion are different from the ones present in preparations without Ca\(^{2+}\) (Fig. 18.8). This indicates that Ca\(^{2+}\) alters the aggregation pathway of S100A6, resulting in homogenous preparations of oligomers versus fibrils. Whereas in the absence of Ca\(^{2+}\) the predominant species are amorphous aggregates and small spherical oligomers (Fig. 18.8A,C, arrowheads and arrows, respectively), in the presence of Ca\(^{2+}\) aggregates are absent and oligomers are larger, probably representing off-pathway species (Fig. 18.8B,D). In fact, the species formed upon Ca\(^{2+}\) incubation are not toxic as assessed by viability assays in cellular models \[14\]. Ca\(^{2+}\) also prevents the generation of fibrils, probably by stabilizing the off-pathway oligomer stage.

**Ex Vivo S100A8/A9 Amyloids Imaged by AFM and TEM**

For the first time, amyloids of pro-inflammatory S100A8/A9 proteins formed in vivo were found in calcified deposits, extracted from the prostate, known as *corpora amylacea* \[12\]. These deposits were classified as local amyloid deposits associated with prostate gland inflammation and aging. Indeed, *corpora amylacea* are frequently observed in the prostate when it undergoes structural remodeling due to age-dependent reduction in reproductive function. This may lead to benign prostatic hyperplasia, observed in 70% of men in their 60s, and even cancer. *Corpora amylacea* inclusions are often adjacent to damaged epithelium and provide a feed-forward loop exacerbating further inflammation, infection and cancerogenesis \[12\]. There is growing evidence that inflammation plays a crucial role in prostate pathogenesis: it is associated with 40–90% of benign prostatic hyperplasia \[70\] and leads to 20% of all human cancers. We have found that protein deposits constitute up to 30% of *corpora amylacea* contents, the remainder being calcified inorganic compounds \[12\]. AFM and TEM analysis of extracts from *corpora amylacea* of seven patients subjected to prostatectomy due to cancer revealed a variety of highly heterogeneous molecular aggregates (Fig. 18.9). These included spherical oligomeric species of ca. 2–3 nm in height in AFM cross-sections (Fig. 18.9A), which closely resembled oligomeric precursors of amyloid fibrils described for numerous peptides and proteins \[57,71\]. In addition, extensive networks of typical fibrillar amyloids 4–8 nm in height and micrometers in length were imaged by AFM (Fig. 18.9B), and straight and rigid fibrils a few hundred nanometers in length were detected by TEM (Fig. 18.9C). There were also larger-scale supramolecular filamentous assemblies, reaching a few micrometers in length and up to ca. 500 nm in thickness (Fig. 18.9D), and large round aggregates of ca. 200–300 nm diameter at half-height and ca. 50–70 nm height in AFM cross-sections (Fig. 18.9E). Comparison of *in vitro* and *ex vivo* amyloids (Fig. 18.5 and 18.9) revealed their general similarity and also their tendency to assemble into large supramolecular complexes.
Macroscopic Amyloid Properties of *Corpora Amylacea* Studied by Optical Microscopy

*Corpora amylacea* inclusions may grow up to millimeter scale, as shown in Figure 18.10A, and may be very heterogeneous in size [12]. In extreme cases, their bulk weight may constitute up to one third of the prostate gland. These inclusions were also characterized by typical amyloid features as observed in optical microscopy. Specifically, they bound the amyloid-specific fluorophore ThT, which presented increased fluorescence under the fluorescence microscope (Fig. 18.10B); the Congo Red dye, also used to identify amyloids, resulted in red staining and Congo Red green birefringence under polarized light, as observed in the optical microscope (Fig.18.10C).

In immunohistochemical analyses, all *corpora amylacea* specimens were stained positively with antibodies recognizing an amyloid conformational epitope [72], S100A8 or S100A9 (Fig. 18.11). This indicates that the S100A8/A9 amyloid material constitutes a significant mass of *corpora amylacea*.
CONCLUSIONS

Above, we have demonstrated that microscopy techniques such as AFM and TEM are able to provide high-resolution insights into the pathways of amyloid assembly of S100 proteins and reveal the complex hierarchy of their amyloid species. S100 proteins are intrinsically amyloidogenic, and we may anticipate that more amyloids of these proteins will be discovered in coming years, both in vitro and in vivo. The direct involvement of pro-inflammatory S100A8/A9 proteins in the formation of corpora amylacea emphasizes the role of inflammation and amyloid formation in age-dependent prostate remodeling and cancerogenesis. As the enhanced level of S100 proteins in general is a characteristic feature of numerous inflammatory conditions, cancers and degenerative conditions taking place in different tissues and organs, they may also effectively contribute to pathology via amyloid depositions. The conformational plasticity of S100 proteins is modulated by Ca$^{2+}$ and other metal ions, and the effect of various environmental factors on their amyloid self-assembly requires thorough investigation. Learning how to regulate the conformational and aggregation properties of these proteins may be a key for effective therapeutic intervention in multiple disorders linked to S100 proteins.

Abbreviations

AFM, atomic force microscopy; ALS, amyotrophic lateral sclerosis; RAGE, receptor for advanced glycation end-products; TEM, transmission electron microscopy; ThT, thioflavin-T; TLR4, toll-like receptor 4.

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