Striatins as plaque molecules of zonulae adhaerentes in simple epithelia, of tessellate junctions in stratified epithelia, of cardiac composite junctions and of various size classes of lateral adherens junctions in cultures of epithelia- and carcinoma-derived cells

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Striatins as plaque molecules of *zonulae adhaerentes* in simple epithelia, of tessellate junctions in stratified epithelia, of cardiac composite junctions and of various size classes of lateral adherens junctions in cultures of epithelia-and carcinoma-derived cells

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**Abstract** Proteins of the striatin family (striatins 1–4; sizes ranging from 90 to 110 kDa on SDS-polyacrylamide gel electrophoresis) are highly homologous in their amino acid sequences but can differ in their cell-type-specific gene expression patterns and biological functions. In various cell types, we have found one, two or three polypeptides of this evolutionarily old and nearly ubiquitous family of proteins known to serve as scaffold proteins for diverse protein complexes. Light and electron microscopic immunolocalization methods have revealed striatins in mammalian cell-cell adherens junctions (AJs). In simple epithelia, we have localized striatins as constitutive components of the plaques of the subapical *zonulae adhaerentes* of cells, including intestinal, glandular, ductal and urothelial cells and hepatocytes. Striatins colocalize with E-cadherin or E–N-cadherin heterodimers and with the plaque proteins α- and β-catenin, p120 and p0071. In some epithelia and carcinomas and in cultured cells derived therefrom, striatins are also seen in lateral AJs. In stratified epithelia and in corresponding squamous cell carcinomas, striatins can be found in plaques of some forms of tessellate junctions. Moreover, striatins are major plaque proteins of composite junctions (CJs; *areae compositae*) in the intercalated disks connecting cardiomyocytes, colocalizing with other CJ molecules, including plectin and ankyrin-G. We discuss the “multimodulator” scaffold roles of striatins in the initiation and regulation of the formation of various complex particles and structures. We propose that striatins are included in the diagnostic candidate list of proteins that, in the CJs of human hearts, can occur in mutated forms in the pathogenesis of hereditary cardiomyopathies, as seen in some types of genetically determined heart damage in boxer dogs.

**Keywords** Adherens junctions · Tessellate junctions · Composite junctions · Intercalated disks · Arrhythmogenic ventricular cardiomyopathy (AC) · Dilated cardiomyopathy (DC)
Introduction

The adhering junctions, i.e. desmosomes (maculae adhaerentes) and adherens junctions (AJs), are major and important elements of the cell-cell connection system in tissues (Farquhar and Palade 1963; for a recent review, see Franke 2009). In the last two decades, the list of the various subtypes of these junctions has been extended by several cell-type-specific forms, including the complexus adhaerens of endothelial cells in certain types of lymph vessels (e.g. Schmelz and Franke 1990, 1993; Schmelz et al. 1994; for a review, see Moll et al. 2009) and the taproot junctions (manubria adhaerentia) of various mesenchymal cells (Wuchter et al. 2007). Of special physiological and medical interest are the myocardiac composite junctions (CJs; areae compositae) that form during the late fetal and postnatal heart development of diverse mammals and that represent densely packed amalgamated arrays of molecules known as desmosomal, peridesmosomal and AJ components of simple epithelia (e.g. Borrmann et al. 2000, 2006; Franke et al. 2006, 2013, 2014; Pieperhoff and Franke 2007; Pieperhoff et al. 2008).

The importance of this type of junction became evident from developmental studies of gene knock-out mice lacking plakoglobin or plakophilin-2 (Bierkamp et al. 1996; Ruiz et al. 1996; Grossmann et al. 2004) and from discoveries of specific, genetically determined cardiomyopathies in human and animal hearts (e.g. Gerull et al. 2004; Antoniades et al. 2006; Heuser et al. 2006; van Tintelen et al. 2006, 2007; Oxford et al. 2007a, b; Posch et al. 2008; Gehmlich et al. 2011; Gaertner et al. 2012; for the recent avalanche of literature, see reviews by Delmar and McKenna 2010; Murray 2012; Rickelt and Pieperhoff 2012; Patel and Green 2014).

In this context, a series of findings concerning hereditary cardiomyopathies in boxer dogs is remarkable; these have been reported to be based on a genetic predisposition for special forms of dilated cardiomyopathy (DC) or arrhythmogenic cardiomyopathy (AC). So far, this seems to be the only pathogenic situation known to involve mutations in a gene encoding a myocardiac member of the striatin family (‘striatin mutations’; e.g. Meurs et al. 1999, 2007, 2010, 2013; Oxford et al. 2007a, 2011). Striatin 1 has repeatedly been reported to be specific for neural cells and functions (e.g. Castets et al. 1996, 2000; Bartoli et al. 1998, 1999; Kachidian et al. 1998; Salin et al. 1998; for other striatins, see also Muro et al. 1995; Moreno et al. 2000; for reviews, see Benoist et al. 2006; Hwang and Pallas 2013). On the other hand, Meurs et al. (2013) have claimed that myocardiac striatin is a desmosomal protein, whereas Breitman et al. (2008) have reported that striatins do not occur in desmosomes but in other kinds of junctions of epithelia and carcinoma cells (for a review, see Hwang and Pallas 2013). The elucidation and examination of possible pathogenic roles of mutated striatins is obviously necessary as these proteins are known as architectonic scaffold molecules able to form oligomers and complexes with other proteins, including kinases and phosphatases, calmodulin and specific Ca\(^{2+}\)-binding proteins, cortactin-binding proteins and signal formation, transduction or vesicle translocation proteins (e.g. Muro et al. 1995; Kachidian et al. 1998; Salin et al. 1998; Bartoli et al. 1999; Moreno et al. 2000; Gaillard et al. 2001, 2006; Yu et al. 2001; Blondeau et al. 2003; Lu et al. 2004; Benoist et al. 2006; Goudreault et al. 2009; Gordon et al. 2011; Bobik 2012; Chen et al. 2012; Tanti et al. 2014; for a review, see Hwang and Pallas 2013).

As the members of the striatin family are highly homologous in their amino acid sequences, and as these isoforms and their splice variants can occur in cell-type-specific patterns, we have decided to address the family of striatins molecules in general in this report and will deal with the diverse cell-type-specific polypeptide isoforms, splice variants and biosynthesis details of striatins in a subsequent protein-chemical-oriented publication.

Materials and methods

Tissues and cell cultures

Bovine tissue samples were obtained from the regional slaughterhouse (Mannheim, Germany) and murine (rat and mouse) tissues were from animals of the laboratory-animal facilities of the German Cancer Research Center (Heidelberg, Germany; for details, see Franke et al. 2006). In addition, tissue specimens from fetal German landrace pigs and 3-year-old boars were obtained from the Institute of Farm Animal Genetics (Friedrich-Loeffler-Institute, Mariensee, Germany; see Rickelt et al. 2011). Cryopreserved human tissue samples, including tumour tissues, were obtained from material taken and examined for diagnostic pathology (Franke et al. 2006; Moll et al. 2009) or were provided by the National Center for Tumor Diseases (NCT, Heidelberg, Germany). In general, the samples were fixed either with 4 % formaldehyde in phosphate-buffered saline (PBS) and embedded in paraffin or were snap-frozen in isopentane that had been precooled in liquid nitrogen and were then stored at −80 °C until use. Protein lysates of frozen tissues were used for SDS-polyacrylamide gel electrophoresis (SDS-PAGE) of peptides (see Franke et al. 2013).

Monolayer cell cultures of various human cell lines were examined, including the breast adenocarcinoma-derived line MCF-7, HaCaT keratinocytes, the colon adenocarcinoma-derived lines CaCo2 and HT29 and the hepatocellular carcinoma-derived cell lines PLC, HepG2, Hep3b and HuH7. Bovine epithelium-derived cell lines included mammary-gland-derived cells of lines BMGE, BMGE+H, BMGE+HE and KE-5. For comparison, rat liver hepatocellular carcinoma cells of the line MH1C1 were studied in parallel.
The non-epithelial cell lines tested included the human cell lines U333/MG, K562, RPMI 8226, HL-60, SV80, WI-38 and RD, the bovine cell line B1, the rat cell line RVFSMC and the mouse cell lines 3 T3 and L929 (for further information, see Boda-Heggemann et al. 2009; Straub et al. 2011; Pieperhoff et al. 2012; Franke et al. 2013). In addition, freshly prepared cultures of human endothelial cells (HUVECs) and rat cardiomyocytes were used as described (cf. Pieperhoff et al. 2012).

Antibodies

Primary monoclonal antibodies (mAbs) and guinea pig polyclonal antibodies (pAbs) were generated against several amino acid sequences (Table 1) of striatin family members obtained as polypeptides synthesized by PSL (Peptide Speciality Laboratories, Heidelberg, Germany). The peptides, coupled via cysteines to keyhole limpet haemocyanin (KLH), were used for the immunization of animals, in particular mice and guinea pigs. Further antibodies against proteins of the striatin family or other molecules used in biochemical and immunolocalization experiments are listed in the Electronic supplementary material (Table S1) and in the publication by Straub et al. (2003).

The protocols in which murine mAbs, guinea pig pAbs and other antibodies were used for immunofluorescence microscopy or for immunoblotting analyses of PAGE-separated polypeptides against AJ molecules or against diverse cytoskeletal proteins were as described elsewhere (Rickelt et al. 2011). The newly generated mAbs and pAbs were routinely compared with “anti-striatin” and “anti-SG2NA” mAbs purchased from Becton-Dickinson (Heidelberg, Germany) or Millipore (Temecula, Calif., USA) and with commercially available polyclonal rabbit antibodies against striatin 4 (“zinedin”; Acris Antibodies, Herford, Germany). Antigen-bound primary Abs were visualized with secondary antibodies coupled to Cy3 (Dianova, Hamburg, Germany) or Alexa 488 (MoBiTec, Göttingen, Germany). For immunoblot analysis, horseradish-peroxidase-conjugated secondary antibodies were applied (Dianova).

Gel electrophoresis and immunoblotting

Protein lysates were analysed by SDS-PAGE, followed by immunoblotting, as described (Rickelt et al. 2011; Pieperhoff et al. 2012; Franke et al. 2013).

Immunofluorescence and immunoelectron microscopy

Methods for immunofluorescence and electron microscopy were as previously described (Franke et al. 2006, 2013; Rickelt et al. 2011; Pieperhoff et al. 2012; Rickelt 2012).

Results

Characterization of striatin proteins and antibodies

At least three genes encoding striatins of highly homologous amino acid sequences (striatins 1, 3, 4) have been identified, each with a series of introns. These genes and introns can result in different cell-type expression patterns of the various isoforms and splice variants. In the present report, we have therefore tried to generate certain polypeptide-sequence-specific antibodies, including some that are specific for certain unique sequence epitopes and others that cross-react between different striatins (see Materials and methods, Table 1).

Using the above antibodies and several that were commercially available, we identified striatins in all normal and tumour cells examined, including single blood cells and tissue cells and in cultured cells and tumour cells (Fig. 1a, b). Whereas some of these antibodies revealed the presence of at least two polypeptide bands of approximately 110 and 100 kDa (Fig. 1a), other sequence-specific antibodies reacted with only one polypeptide (cf. Fig. 1b, b'). The common bands identified by some of the antibodies often appeared rather faint on some tissues, notably those of liver and heart, but were much more intense at higher protein loads or after extended immunoblot exposure times (see also Electronic supplementary material, Fig. S1). When various preparations of mammalian heart tissue or murine cardiomyocyte cell cultures were compared, a band with an Mₚ of approximately 110 kDa was always seen and, in some preparations, was accompanied by a (mostly minor) band of a lower Mₚ (the obvious difference with respect to the SDS-PAGE immunoblot data of Castets et al. 2000, who reported only cardiac...
polypeptides of lower Mr values, i.e. approximately 94 and 100 kDa, cannot yet be explained).

Colocalization experiments

Because of the dense-packing of cytoskeletal and cell junction components, special and carefully controlled antibody binding and differential washing protocols are needed to distinguish true and specific epitope binding from the various forms of structure and protein “stickiness”. In the present study, we have generally included diverse washing steps to remove non-specifically bound (“sticky”) material from the structures in question, and in a series of cases, this required brief (5, 10 or 15 min) rinsing with mild detergent-containing buffers and/or acetone solutions. To illustrate the importance of such differential washing steps, we include here, as an example, the binding and release reactions of desmoplakin and plectin (Electronic Supplementary Material, Fig.S2). Whereas desmoplakin is known as an extraction-resistant, intensely binding component of the CJs in the intercalated disks of the myocardium, plectin is, in addition, known for its marked “sticky” behaviour, i.e. binding that is not immunologically determined. Consequently, the plectin reaction with the sarcomeric Z-lines can (and should) be removed by differential washing (Fig.S2; cf. Fig.S2a, b). On the other hand, in the course of these washing steps, a significant portion of the plectin antibodies remain bound to the CJs, thus resulting, together with established CJ markers, such as desmoplakin, in a typical yellow merged reaction colour (Fig.S2a, b’). For example, the differential localization reaction of a “sticky” protein and a CJ-specific plaque protein is shown in Fig. S3, comparing striatin as a CJ-specific protein with α-actinin as a “sticky” sarcomeric Z-line protein (Fig. S3; cf. Bennett et al. 2006). By contrast, various types of antibodies against α-actinin colocalize with high precision and intensity (see the yellow merged pattern in Fig. S4). Consequently, extensive differential washing treatments have been included in the immunolocalization experiments of this study.

**Fig. 1** Results of SDS-polyacrylamide gel electrophoresis (SDS-PAGE)-separated polypeptides as obtained by immunoblot reactions (a, b, b’, c’). The antibodies used were monoclonal antibody (mAb) “Striatin” (Becton-Dickinson; a, b, c’) and polyclonal antibody (pAb) raised in guinea pig (gp), namely “striatin mix” of NTB, 268B, 301B and CTB (b’). The tissue and cell lysis protein preparations used were from human heart (lane 1), tongue mucosa (lane 2), liver tissue (lane 3) and the following human cell culture lines: PLC (lane 4), HaCaT (lane 5), SV80 (lane 6), A498 (lane 7), CaCo2 (lane 8), A431 (lane 9), HeLa (lane 10), HUVEC cells (secondary cell culture; lane 11), K562 culture 01 (lane 12), K562 culture 02 (lane 13) and RPMI 8226 culture 01 (lane 14). In the results shown in c and c’, the following materials were used: human heart tissue (lane 1), bovine heart tissue (lane 2), murine heart tissue (mouse; lane 3), murine HL-1 culture line of cardiomyocytes (lane 4), and a primary cell culture of neonatal rat cardiomyocytes (lane 5). Note the dominant immunoblot polypeptide band at approximately 110 kDa in all tissues and cell cultures, except for the weak reaction in lane 1 of a, which is, however, more noticeable at higher protein loadings. In addition, an immunoreactive band at approximately 100 kDa is seen in most lanes of a, b and in lanes 2, 4 of c’. In specific lanes of a and in lane 8 of b, additional bands are notable that have not yet been characterized. Further reaction bands are seen at approximately 100 kDa (a, b, lanes 2, 4 in c) and at approximately 142 kDa (lanes 4, 5 in a, lane 8 in b).
Light microscopic immunolocalization in simple epithelial tissues

Using various antibodies against members of the striatin family on cryostat sections of diverse forms of simple epithelia, we have obtained a distinct immunolocalization pattern marking the upper portion in the *zonula adhaerens*. Figure 2a-a′′, for example, presents bovine intestinal epithelium. At the limited resolution in such semithin sections, the striatin reaction, for the most part, overlaps with that of the apical-most desmosomes (yellow merger colour), whereas the desmosomes lying more basally on the lateral cell membranes do not react at all or overlap optically with striatin antibodies (e.g. Fig. 2b). In thin sections, however, one can even locally often distinguish a specific thin apical striatin-positive *zonula*-like structure from the subjacent general *zonula adhaerens* region reacting, for example, with α-catenin, β-catenin, p120, p0071 and protein ZO-1 (for β-catenin, see Fig. 2c–e). In such thin sections, the striatin and β-cadherin-positive zones can also be distinguished from the adjacent occludin- and claudin-positive *zonulae occludentes* of tight junctions (TJ; not shown). This subapical *zonula* region is also different from the *zonula* reaction sites of the 21-kDa transmembrane protein PERP (Fig. 2′′); the apical zone under discussion here is demarcated by the bracket symbol in Fig. 2b). Similar results have been obtained in diverse tissues with adluminal simple epithelia, including salivary and other glandular epithelia, duct epithelia and pulmonary epithelium, bladder urothelium and the seminiferous and excurrent duct epithelia of the testis (cf. Domke et al. 2014). In bovine muzzle epithelial glands and ducts (Fig. 3), the marked striatin *zonula* immunostaining is seen in both the secretory and the ductal cells. Essentially identical results have been obtained in all five mammalian species examined.

As the polar organization of the hepatocytes in mammalian liver tissue represents an especially complex junction, and as liver physiology and diseases are of special importance, we have performed detailed double- and triple-label high-resolution immunolocalization reactions on cryostat liver sections of the five mammalian species used, namely mouse, rat, pig, cattle and human. Figure 4 presents the results obtained by double-label immunofluorescence microscopy, comparing the punctate, rather regularly spaced desmoplakin reaction sites along the bile canaliculi with the thin and distinct, but also intensive, striatin reaction of the apical *zonula adhaerens* structures (Fig. 4a-a′, b-b′). Moreover in cross-sections through the bile canaliculi, we could demonstrate (Fig. 4c–c′) the entire subapical plasma membrane reaction of striatin in direct comparison with the surrounding desmosomes. These results were identical in all five species and were also similar to those obtained for other *zonula adhaerens* markers, including the proteins myozap (Rickelt et al. 2011) and LUMA (Franke et al. 2014).

Light microscopic immunolocalization in stratified epithelia

Epithelia of this category are characterized by variously sized and variously structured interdesmosomal regions that can be studded not only with “gap junctions”, but also with single molecules or “islands” of TJ and/or AJ molecules (“tessellate junctions”); cf. Franke and Pape 2012; Franke et al. 2013. The patterns of these interdesmosomal cell-cell junction structures vary markedly not only between the various types of epithelia, but also in the various cell layers.

Punctate and *fascia*-like striatin immunolocalization reactions have also been noted in the various stratified epithelia examined. For example, the distribution of striatin-containing portions in tessellate junction layers of bovine tongue mucosa is shown in comparison with immunostaining for β-catenin in Fig. 5. In the interdesmosomal cell-cell contact regions of these stratified tissues, small punctate *fascia*-like or even more extended striatin reaction sites are often seen, mostly showing colocalization of AJ molecules with TJ markers such as occludin, because of spatial overlap. Moreover, in several stratified epithelia, striatin immunostaining is not restricted to colocalization areas with other AJ proteins but has selectively been noted in upper layers, positionally equivalent to the upper *stratum spinosum* and the *granulosum* layers of the epidermis, even in regions that appear totally negative for proteins including the catenins, p120, p0071 and protein ZO-1 (see Fig. 5, upper portion). The reactive structures and the intensities of the various members of the striatin protein family can differ in the different stratified epithelia, i.e. epidermis, oral and lingual mucosa regions, oesophagus, pharynx epithelium and stratified thymic reticulum epithelium (“Hassall bodies”). Therefore, we have decided to devote a special future article to the complex patterns of AJ protein localizations in the distinct substructures of tessellate junctions of mammalian stratified epithelia and in tumours and cell cultures derived therefrom.

Immunolocalization in myocardiac tissues

In view of the molecular architectonic, functional and medical importance of CJs in the intercalated disks, and in view of the special roles of such molecules in the pathogenesis of a series of heart diseases and “sudden death” forms, we have carefully examined and compared the five mammalian species mentioned by immunofluorescence microscopy. Moreover, as a single striatin polypeptide appeared to be the predominant, if not exclusive isoform in myocardiac cells in situ (see Fig. 1a, c′), we made certain, in all cases, that the antibodies specific for this striatin were included in the experiments.

As shown for the example of boar heart (Fig. 6), striatin is highly enriched in the CJs, usually showing colocalization with N-cadherin and β-catenin (Fig. 6a-a′′, b-b′′) and with p120, desmoplakin, plakophilin-2 and desmoglein-2 (Dsg2;
not shown). Essentially identical colocalization results have been obtained for bovine cardiomyocytes in situ (Fig. 7a–a’’ presents, for example, colocalization with desmoplakin). Several other AJ plaque proteins such as ZO-1 also colocalize with striatin in CJs but here striatin label has not been detected in the zonulae of the interspersed blood capillaries (Fig. 7b, b’).

Colocalization of striatin with other CJ molecules has also been found in a significant proportion of the junctions of the Purkinje network of conducting cells (Fig. 7c–c’’; for an extensive recent review of cardiac conduction cells, see Mezzano et al. 2014). In addition, we have noted, in the conductive cells, a few striatin-positive junctional reaction sites that are negative for all desmosome-specific marker proteins and other junctions that are desmoplakin- and plakophilin-2-positive but negative for all striatin antibodies tested (not shown).

**Fig. 2** Immunolocalization of striatin in the zonula adhaerens of bovine intestinal epithelial cells. a–a’’ Double-label confocal laser scanning immunofluorescence microscopy, showing striatin (a, red; mAb mouse [m]) in a relatively narrow apical zone in partial colour reaction overlap with desmoplakin (a’, green; pAb gp). This double-label immunoreaction is seen in a” and on a phase contrast background in a’”. Note, however, that striatin is restricted to the subapical ring (zonula adhaerens; L, lumen), whereas desmoplakin is also located in the numerous desmosomes of the basolateral cell-cell contacts. b Higher resolution micrograph showing the distinct separation of the striatin-positive zonula (red) and the basolateral desmosomes (green). c–e Differential immunostaining reactions of striatin (red; mAb m) and β-catenin (green; rabbit [rb] antibody), indicative of zones of colocalization (yellow) and local segregation (red). f–f’’ Double-label immunofluorescence microscopy showing the same tissue after localization of proteins PERP (f, f’, f’’, red; mAb m) and striatin (f–f’, green; gp). Note that striatin is restricted to a thin upper line of the zonula adhaerens (green), whereas protein PERP is seen in a slightly lower zone and in special larger punctate structures both at the zonula and at the basolateral membranes (bracket and bottom in f, respectively). Bars 20 μm (a, f), 5 μm (b–e).

**Fig. 3** Double-label confocal laser-scanning immunofluorescence microscopy of cryostat sections through a bovine muzzle epidermis region rich in glandular and ductal epithelium. a Survey of cross- or obliquely-sectioned gland structures showing the general frequency of desmosomes (green; desmoplakin, gp antibody) and zonulae adhaerentes positive for striatin (red; mAb m). a’ Higher magnification illustrating the differential localization of these two structures. Bars 20 μm.
In both murine species examined, the colocalization of striatin with β-catenin (Fig. 8a—a′'), p120 and p0071 and with desmoglein-2, desmoplakin, plakophilin-2 and plakoglobin have also been observed. However, in a number of experiments, we have also detected small (“dot-like”) cytoplasmic reaction sites that are positive only for striatin or only for specific CJ partner molecules such as β-catenin (see Electronic supplementary material, Fig. S5), p120 and p0071 or α-catenin (not shown).

Unsurprisingly, the same kind of results were also obtained in our extensive localization studies of human myocardium. As shown in cross-sections of intercalated disks (Fig. 8a—a′′, c)
and in grazing-horizontal sections of intercalated disks (Fig. 8b–b’’’), pronounced colocalization was typical for at least one isoform of striatins with N-cadherin (Fig. 8a–a’’’), β-catenin, p120 and p0071 (not shown) and with desmoplakin (Fig. 8c), desmoglein Dsg2, plakophilin-2 and plakoglobin (not shown). In special control experiments, we also used two different types of striatin antibodies, namely, one that cross-reacted with various striatins and one that reacted only with the major cardiac striatin polypeptide. Again, near-complete colocalization was observed (Fig. 8d).

Colocalization of striatin(s) in the plaques of CJs has also been found for plectin, an extremely large protein previously described in association with various other contractile and cytoskeletal proteins (Wiche et al. 1983; Wiche 1989; Andrä et al. 1997; for biochemical data, see also Wiche et al. 1982; Pieperhoff et al. 2012) and

**Fig. 5** Double-label confocal laser scanning immunofluorescence microscopy showing one of the subforms of cell-cell tessellate junctions in a multistratified epithelium, namely the ventral part of bovine tongue mucosa. a–a’’ Striatin (red; mAb m) demonstration in regional substructures, including polar or fascia-like tessellate junctions; the β-catenin-positive portion (green; pAb gp) of the tessellate junctions extends over much larger cell-cell contact areas. Bar 20 μm
ankyrin-G (Electronic supplementary material, Fig. S6), confirming the data of Makara et al. (2014; see also Mohler et al. 2004; Sato et al. 2011; for a review, see Bennett and Healy 2009). To demonstrate the specificity and intensity of the binding of ankyrin-G and plectin to other CJ plaque proteins and, notably, also to the protein myozap (see also Pieperhoff et al. 2012) and striatin (Fig. S6f), gradual “buffer wash treatments” of cryostat sections have been regularly performed (see also previous sections).

Light microscopic immunolocalization of striatin in cultured epithelial and myocardial cells

Localization studies of striatin with α- and β-catenin, with other AJ markers and with desmosomal molecules...
have also been performed on cell culture monolayers, including epithelium or carcinoma-derived cells and cardiomyocyte-derived cells (for biochemical demonstrations of the presence of striatins in such cells, see Fig. 1a, b and Electronic supplementary material, Fig. S1; for immunofluorescence microscopy, see Electronic supplementary material, Figs. S7–S10). Striatins have been identified as major components not only in the cell-cell connecting \textit{zonulae adherentes} of primary cultures. Figure S7, for example, shows a monolayer culture of rat myocardiac cells taken 2 days after birth (cf. Pieperhoff et al. 2012). This micrograph...
also demonstrates the absence of striatin in certain desmosome-related structures, including the variously sized intracellular assemblies of desmoplakin-rich material. Striatin has also been seen in continually-appearing cell-cell contact AJs of all epithelial cell culture monolayers examined (Figs. S8–10 present examples of the human breast carcinoma line MCF-7). Thus, striatins have to be counted among the obligatory constituents of AJs in tissues and in cell cultures. As striatins also occur in single cells in culture and in the living mammalian body (see Fig. 1 and Electronic supplementary material, Figs. S6–S9), one has to conclude that the synthesis and stability of striatin(s) are not dependent on established cell-cell junctions.

Remarkably, the integration of striatin(s) into the AJ zona Adhaerens plaque structures is not restricted to completed assembly at the cell-cell contacts but can be detected in small puncta- or fascia-like structures in the cytoplasm, even in juxtanuclear regions, or in short plasma membrane intercepts before the formation of a continuous zona adhaerens (Figs. S8, S9). In such situations, we find it especially surprising that even the
newly formed AJ molecules are often closely associated with TJ proteins such as claudins and/or occludin (Figs. S8–S10).

Immunoelectron microscopic localizations of striatins

Using snap-frozen tissues or monolayer cell cultures, we have been able to localize specific striatins on the plaques of cell contact regions of the AJ type, as is shown for bovine liver tissue in Fig. 9a, b. Again, desmosomes or other categories of junctions (gap junctions, TJs) are not immunogold-labelled at all (e.g. the junction labelled D in Fig. 9c). Moreover, the immunogold reaction sites are all associated with cytoplasmic plaques of AJ structures. Particularly eye-catching in this tissue is the zonula adhaerens labelling extending over the subapical junction region bordering on the bile canaliculi (BC in...
Fig. 9d), whereas no striatin label is associated with the apical villus-like cytoplasmic processes extending into the bile canalicular interior.

Even more extended and often dense immunolocalization reaction products have been seen in the plaques of interdesmosomal AJ-type cell-cell contact regions of densely grown monolayers of human breast carcinoma MCF-7 cells (Fig. 9e, f), hepatocellular carcinoma PLC and colon carcinoma CaCo2 cells (not shown) and in human epidermis-derived HaCaT tumour cells (not shown). Here, large amounts of antibody-bound heavy metal grains are exclusively concentrated on the submembranous AJ plaques (see Fig. 9f), whereas neither gap junctions nor TJs show marked enrichment.

In stratified epithelia, immunoelectron reaction is also restricted to interdesmosomal regions. Figure 10a-h shows striatin labelling by gold-silver grains at such regions (desmosomes are numbered in Fig. 10a) with only a few small immunogold grains. By contrast, Fig. 10b presents clusters of larger metal label grains in positions that, in some cases, might be equivalent to local puncta adhaerentia within a tessellate junction (regions denoted by brackets in
Table 2 Constitutive molecules of composite junctions (areae compositae) in the intercalated disks of mammalian cardiomyocytes

| Transmembrane cadherins          | Plaque proteins          |
|------------------------|--------------------------|
| **Adherens junction molecules** | **Adherens junction molecules** |
| N-cadherin            | α-Catenin (αE + αT)      |
| Cadherin-11           | β-Catenin                |
|                       | Protein p120             |
|                       | Protein p0071            |
|                       | Protein ARVCF            |
|                       | Plakoglobin              |
|                       | Myozap                   |
|                       | Striatin(s)              |
|                       | Protein ZO-1             |
| **Desmosomal molecules**   | **Desmosomal plaque molecules** |
| Desmoglein 2 (Dsg2)      | Plakoglobin              |
| Desmocollin 2 (Dsc2)     | Plakophilin 2 (Pkp2)     |
| **Other molecules**       | Desmoplakin              |
| Plectin                |                          |
| Ankyrin-G              |                          |
| Protein PERP           |                          |
| Protein LUMA           |                          |

†Armadillo repeats-containing proteins are underlined

Fig. 10b). An even more extended and intensive interdesmosomal striatin labelling pattern is seen in Fig. 10c). In general, the sizes and the relative positions of striatin immuno- reaction products vary (for light microscopic comparisons, see Electronic supplementary material, Fig. S5). In Fig. 10d, the label extends over the entire interdesmosomal region, whereas the antibody-linked heavy metal grains in Fig. 10e-h show a higher tendency to cluster in regions near desmosome margins. In view of the frequent close vicinity of striatin localization sites with desmosomal margins (see above), we have examined the proteins of desmosomal fractions from calf muzzle epidermis, tongue mucosa and oesophagus tissue prepared in the Heidelberg laboratory (see Mueller and Franke 1983; Kapprell et al. 1985) by SDS-PAGE and immunoblotting, with and without immunoprecipitation and protein cross-linking experiments (cf. Straub et al. 2011; Pieperhoff et al. 2012) but we have found no indications of the occurrence of striatins within desmosomal structures.

Discussion

Members of the striatin family of proteins are by no means specific for, or especially abundant in, neuronal or other cells of the nervous system as originally reported (for references see Introduction). As shown in this study, they are synthesized and integrated into protein complexes, dispersed particles or plaques of cell junctions of most, if not all mammalian cells, in single cells and in cultured cells and cells of tissues (see also Hwang and Pallas 2013).

For the sake of clarity, we consider it important to emphasize that our results show that the classification of members of the striatin family as “desmosomal proteins” (Meurs et al. 2013) is not correct. We have not detected striatin(s) in any type of desmosome or desmosome-related structure. On the contrary, striatins are exclusive and constitutive components of AJs such as the *zonulae adherentes* of polar epithelia, of certain punctate or fascia-like substructures of the desmosomal complexes in stratified epithelia, and of the CJs in the intercalated disks of mammalian heart (for the latter localization in boxer dog, see also Meurs et al. 2010). Consequently, striatins (certainly the major cardiac isoform) should be added to the diagnostic “control list” of markers for hereditary AC and DC damage (Table 2; cf. Asimaki et al. 2009).

Our results also clearly show that striatins are not components of TJs, although they can often be seen in the vicinity of TJs. The most convincing argument for this conclusion is presented by the CJs of the myocardial intercalated disks from which TJ-specific molecules and structures are totally absent. Similarly convincing examples are provided by several cultured cells with large regions positive for striatins and other AJ molecules but negative for TJ markers.

Additional locations of members of this protein family might occur in which the striatins as scaffold proteins are masked by specific complex partner molecules. Indeed, the members of this protein family are typical scaffold proteins able to form not only oligomeric complexes, but also complex “multimodular” structures, including some involved in diverse signalling and regulatory functions (see Muro et al. 1995; Castets et al. 1996, 2000; Bartoli et al. 1998, 1999; Kachidian et al. 1998; Salin et al. 1998; Moreno et al. 2000, 2001; Baillat et al. 2001; Gaillard et al. 2001; Yu et al. 2001; Lu et al. 2004; Joshi-Mukherjee et al. 2008; Gordon et al. 2011; Chen et al. 2012; for a recent general review on scaffold proteins see Garbett and Bretsch 2014). Consequently, we now need to examine the possible occurrence of such striatin-binding proteins and striatin-typical functions in plaque complexes of, for example, epithelial AJs and myocardial CJs. Similarly important are detailed experimental analyses, including gene knock-out studies, to determine which of the so highly related and so similarly sized striatin forms are involved in certain pathogenic conditions, such as the aforementioned cardiac AC and DC damage.

Whether the pathogenic effects of certain CJ molecule mutations on cardiomyopathies such as ACs or Brugada syndrome take place in junctions between cardiomyocytes or in any of the special junctions connecting the conductive Purkinje fibre cells (for details, see Pieperhoff et al. 2010; Mezzano et al. 2014) remains unclear. Also unknown is whether these effects are direct or indirect, e.g. by involving...
connexin43 or connexin40 of the intimately associated gap junctions or the Nav1.5 sodium ion channels, and decisive experimental results are needed. Such indirect reactions via other adjacent cell-cell contact structures have repeatedly been discussed by a series of authors (Oxford et al. 2007a, b, 2011; Sato et al. 2009, 2011; Cerrone et al. 2012, 2014; Delmar and Liang 2012; Gomes et al. 2012; Agullo-Pascual et al. 2013, 2014; Meens et al. 2013; Noorman et al. 2013; Cerrone and Delmar 2014; Lyon et al. 2014; Vreeker et al. 2014; for earlier references and general reviews, see also Saffitz 2007, 2011; Delmar and McKenna 2010; Delmar and Makita 2012; Murray 2012; Rickelt and Pieperhoff 2012; Rizzo et al. 2012a, b). Finally, we cannot yet exclude that both direct and indirect effects of the mutated or otherwise modified molecules contribute to the pathogenetic effects mentioned. One hope is that detailed studies of the animal pathogenesis examples and transgenic experimental possibilities (for some related references, see Table 3 in Rickelt and Pieperhoff 2012; see also Fox et al. 2007; Vatta et al. 2007) will help in elucidating the cardiomyopathic mechanisms involved.

The recognition of striatin as a multimodular scaffolding protein that occurs in the cytoplasmic plaques of various kinds of cell junctions has also to be discussed in comparison with other scaffolding plaque proteins such as protein ZO-1 of various tight and adherens junctions (AJs), as well as AJ α-catenin and the ERM protein-binding component EB50 found at the actin microfilament associations with microvillar membranes (for a review, see Garbett and Bretscher 2014). Such scaffolding proteins can also form a diversity of other complexes located in other structures or regions in which they might be masked for certain cell-type-specific antibodies but do indeed bind to antibodies reactive with other epitopes on the same molecule. As a result, the same scaffolding protein might immunocytochemically appear positive in one structure but negative in another. Consequently, such selective masking of one conformational domain might result in completely negative immunolocalization sites of the same protein that is positive with other antibodies (see, for example, the results obtained for protein LUMA in the report of Franke et al. 2014).

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