The presence of iduronic acid in chondroitin/dermatan sulfate changes the properties of the polysaccharides because it generates a more flexible chain with increased binding potentials. Iduronic acid in chondroitin/dermatan sulfate influences multiple cellular properties, such as migration, proliferation, differentiation, angiogenesis and the regulation of cytokine/growth factor activities. Under pathological conditions such as wound healing, inflammation and cancer, iduronic acid has diverse regulatory functions. Iduronic acid is formed by two epimerases (i.e. dermatan sulfate epimerase 1 and 2) that have different tissue distribution and properties. The role of iduronic acid in chondroitin/dermatan sulfate is highlighted by the vast changes in connective tissue features in patients with a new type of Ehler–Danlos syndrome: adducted thumb-clubfoot syndrome. Future research aims to understand the roles of the two epimerases and their interplay with the sulfotransferases involved in chondroitin sulfate/dermatan sulfate biosynthesis. Furthermore, a better definition of chondroitin/dermatan sulfate functions using different knockout models is needed. In this review, we focus on the two enzymes responsible for iduronic acid formation, as well as the role of iduronic acid in health and disease.

Introduction

Dermatan sulfate (DS) is a glycosaminoglycan (GAG) that is distinguished from chondroitin sulfate (CS) by the presence of iduronic acid (IdoA), the C-5 epimer of D-glucuronic acid (GlcA). IdoA occurs in variable proportions in DS (Fig. 1A) and, as a result of the different position of the carboxyl moiety (Fig. 1B), it generates a more flexible polysaccharide chain, allowing specific interactions with several proteins and polysaccharides. To form CS/DS, three specific enzymes, dermatan sulfate epimerase 1 (DS-epi1), dermatan sulfate epimerase 2 (DS-epi2) and dermatan 4-O-sulfotransferase 1 (D4ST1), are required [1]. These enzymes are differently organized in various tissues and, under different physiological conditions, they generate CS/DS of a very different structure. DS is found relatively late in the evolutionary tree and first appears in molluscs, sea urchins and sea cucumbers. It is then found in ascidians and in the whole vertebrate phyla [2]. However, it is absent in Caenorhabditis elegans and Drosophila melanogaster. The present review presents the structure, function and biosynthesis of these structurally different CS/DS polymers and explains how
they are modified in response to different physiological and pathological processes.

**Structure of CS/DS**

CS/DS chains are found on at least 32 different core proteins forming proteoglycans (Table 1). Six of these are also substituted with heparan sulfate. Some of these proteoglycans, such as CD44, α5β1 integrin and collagen XV, are only part-time proteoglycans.

CS is a long polysaccharide consisting of the repeating disaccharide units GlcA and N-acetyl-galactosamine (GalNAc), attached to serine residues of core proteins. The chains from eukaryotic organisms are extensively modified by sulfation, yielding six different disaccharides: GlcA-GalNAc residues (O unit), GlcA-GalNAc-4-sulfate (A unit), GlcA-GalNAc-6-sulfate (C unit), GlcA-GalNAc-4,6-disulfated (E unit). The GlcA residue can also be sulfated at the 2-position giving rise to B units (GlcA-2-sulfated-GalNAc-4-sulfated) and D units (GlcA-2-sulfated-GalNAc-6-sulfated) [3]. Even more complex sulfation patterns have been described in the invertebrate phyla [2].

An important modification is the epimerization of GlcA residues to IdoA residues by C-5 inversion at the polymer level of a (β-GlcA-1,3-β-GalNAc-1,4)n substrate (Fig. 1B) [4]. Individual saccharide units in CS/DS can exist in different conformations depending on their structural arrangement. IdoA residues allow flexibility given their ability to switch between 1C4 (chair), 2S0 (skew boat) and 4C1 (chair) conformations (Fig. 1C), whereas GlcA residues are less flexible and exist in the 4C1 (chair) conformation [5]. IdoA can occur in three different arrangements: (a) as a single IdoA-containing disaccharide surrounded by GlcA containing disaccharides; (b) in structures where they alternate with GlcA containing disaccharides or (c) in long blocks of adjacent IdoA-containing disaccharides (Fig. 1A). The sulfation pattern differs according to the IdoA distribution because IdoA blocks are mainly

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**Fig. 1.** Structure of CS/DS and conformations of IdoA. (A) The domains of variable length containing blocks of IdoA, alternating IdoA and GlcA or blocks of GlcA. (B) The epimerase reaction. (C) Conformations of IdoA.
Table 1. CS/DS PGs and functions of the CS/DS chain. NA, not analyzed.

| PG               | Presence of IdoA | Functions of PG                                                                 | CS/DS binding proteins and CS/DS functions |
|------------------|------------------|---------------------------------------------------------------------------------|--------------------------------------------|
| **Extracellular matrix** |                  |                                                                                 |                                            |
| Aggrecan         | NA               | Chondrocyte differentiation, motility, proliferation and metastasis [79,80]; ECM assembly [81] | Water retention                            |
| Versican         | IdoA+            | Increases differentiation, motility, proliferation and metastasis [79,80]; ECM assembly [81]; TGF-β interaction [82]; self-association [83]; modulation of proliferation, survival, migration and angiogenesis [84]; coagulation [60]; LDL interaction [54]; Borrelia invasion [65]; α-defensin targeting [66]; progeroid and Ehlers–Danlos syndromes [85] | FGF family, L- and P-selectin, chemokines |
| Decorin          | IdoA+            | TGF-β interaction [82], self-association [83]; TGF-β interaction [82]; self-association [83]; modulation of proliferation, survival, migration and angiogenesis [84]; coagulation [60]; LDL interaction [54]; Borrelia invasion [65]; α-defensin targeting [66]; progeroid and Ehlers–Danlos syndromes [85] | FGF2, FGF7, HGF, HCII, αβ1 integrin, tenascin-X, fibril formation, DS:DS self-association [86] |
| Biglycan         | IdoA+            | Interactions with TGF-β [87]; BMP4/chordin [88]; collagen I [89]; associated with tumour in gastric tissue [90] and endothelial cells [91]; involved in inflammation and development [92,93]; neuronal survival [94]; bone development and osteoporosis [95,96] | HCII, FGF family                           |
| Epiphycan        | IdoA+            | Chondrocyte differentiation [97] and matrix organization in the growth plate [98] | NA                                         |
| Collagen IX      | NA               | Organization of cartilage [99], associated with fibroblasts in colon cancer | NA                                         |
| Collagen XII     | NA               | Organization of cartilage and skin [100]                                       | NA                                         |
| Collagen XIV     | NA               | Organization of cartilage and skin [101,102]                                     | NA                                         |
| **Cell surface** |                  |                                                                                 |                                            |
| Betaglycan       | NA               | TGF-β presentation [103,104] and suppression of cancer progression and metastasis [105]; binds inhibin and suppresses activin signalling [106] | NA                                         |
| Syndecan-1       | IdoA+            | Regulation of tumour cell survival and proliferation, growth factor and cytokine binding, adhesion [107–109] | Midkine, pleiotropin, FGF                   |
| Syndecan-3       | NA               | Role in human labour [110,111], adhesion, growth factors co-receptor, neurite outgrowth [112], expressed in tumour stromal vessels [113] | NA                                         |
| Syndecan-4       | NA               | Interaction with Frizzled7 and Dishevelled, regulates noncanonical Wnt signalling and convergent extension movements in Xenopus [114], regulates neural crest cells migration [115] and neural induction via extracellular signal-regulated kinase and protein kinase C pathways [116], adhesion, growth factors co-receptor [109], wound healing and angiogenesis [117], up-regulated in cancer and mediator of cell spreading [118] | Midkine, pleiotropin, bFGF [109] |
| CD44             | IdoA+            | Tumour growth, angiogenesis, metastasis, migration, HGF binding [119]           | Migration, HGF                             |
| NG2              | NA               | Regulates tumour cell growth, motility and survival [120]                        | Differentiation, proliferation and motility, PDGF-AA and FGF2, adhesion [121] |
| α5β1 integrin    | NA               | Fibronectin binding, regulation of adhesion and migration [122]                 | NA                                         |
| **Nervous system** |                  |                                                                                 |                                            |
| Neuropilin-1     | NA               | Metastasis, neuronal guidance, regulation of cell migration [123]                | VEGF signalling                            |
| Neurocan         | NA               | Up-regulated in astrocytoma [124], neurite outgrowth, growth factors binding, brain ECM organization [125] | N-CAM, HB-GAM, amphoterin                   |
4-sulfated with some adjacent sulfated IdoA residues (iB) close to the nonreducing terminal of the blocks [6,7]. The short GlcA blocks are mostly 4-sulfated, whereas longer blocks also contain 6-sulfated GalNac residues [8]. The resulting CS/DS chains therefore contain different domains that enrich their functional properties. The presence of alternating IdoA-GlcA or isolated IdoA has been overlooked in many cases. Furthermore, the content of IdoA varies within the same proteoglycan depending on the tissue of expression [6] and physiological conditions [9]. This is the case for decorin, which is highly iduronated in skin. In bone decorin, however, IdoA is virtually absent [6]. Given the fact that a chain containing IdoA always contains GlcA, the name CS/DS indicates the hybrid nature of the chain.

The structural characterization of CS/DS takes advantage of specific lyases such as chondroitinase ABC, AC and B, which specifically degrade galactosaminoglycans depending on the presence of IdoA or GlcA. The development of high-resolution HPLC systems with pre- or post-column fluorescent derivatization has enabled the separation and quantitation of the various building blocks [10,11]. These methods can only determine the degree of sulfation and the occurrence of IdoA- and GlcA-blocks. However, detailed sequence analysis is not possible. The advent of sensitive MS with different fragmentation procedures has lead to promising results [12,13]. Recently, the complete sequence determination of the chondroitin sulfate in bikunin has been accomplished [14].

### Biosynthesis of DS

DS-epi1 and DS-epi2 catalyze the formation of IdoA, the stereoisomeric form of GlcA, by repositioning the C5 carboxyl group in space (Fig. 1B). DS-epi1 (coded by the gene DSE) and DS-epi2 (coded by the gene DSEL) are both ubiquitously expressed and have common structural features [15,16].
DS-epi1 and 2 share a common N-terminal epimerase domain (Fig. 2A) with 51% amino acid sequence identity between the two enzymes. The secondary and tertiary structures of this domain in the two enzymes are very similar. DS-epi1 has a C-terminal domain of unknown function and three-dimensional structure. There is a similarly positioned domain in DS-epi2 with unknown function and structure. These two domains in the two epimerases do not have significant homology. In addition, in DS-epi2, there is a C-terminal domain, which has 16% amino acid identity with chondroitin-O-sulfotransferase 1, recognized in the database as a CS/DS-O-sulfotransferase domain (Fig. 2A), suggesting that DS-epi2 is an enzyme with dual epimerase and O-sulfotransferase activity. Other enzymes for GAG biosynthesis have been shown to accommodate dual activities [17,18]. The functional epimerase domain of the DS epimerases comprises two structural domains: one mainly composed of \( \alpha \)-helices and one of \( \beta \)-sheets (Fig. 2B). These two domains of DS-epi1 were modelled on the crystal structure of heparinase II [19]. At their boundary, they form a groove, where the substrate is positioned. Some amino acids that are essential for enzyme activity have been identified and a catalytic mechanism has been proposed. Histidine 450 abstracts the C5 proton from one side of the sugar plane of GlcA. This is followed by cleavage or glycosidic linkage between GalNAc and GlcA to generate a C4–C5 double bond containing hexuronic acid intermediate. This structure is finally protonated by histidine 205 adding a hydrogen at the side of the sugar plane that is opposite to the abstraction side.

**Fig. 2.** (A) DS-epi1 and DS-epi2 domain structures. (B) Three-dimensional modelling of the DS-epi1 epimerase domain based on the crystal structure of heparinase II. A chondroitin sulfate tetrasaccharide is positioned in the groove containing the active site.
Finally, the glycosidic link is recreated. As a result of the reaction, the carboxyl group has a different spatial orientation in the IdoA epimer than in the starting GlcA. A prerequisite for activity is the presence of at least three of the four N-glycans.

DSE and DSEL are on chromosomes 6 and 18, respectively [15,20]. The exon/intron organization of the two enzymes is very different because DSE has six exons and the coding sequence spans five exons, whereas DSEL has only two exons (being the whole ORF present in the single exon 2).

The epimerase activity is highly expressed in the spleen, stomach, uterus, ovary, kidney and lung. In the brain, the activity is low and no activity is found in serum [21]. By analyzing the total activity in tissues and mouse embryonic fibroblasts of DS-epi1−/− and DS-epi2−/− mice, it is possible to show that DS-epi1 is the predominant epimerase in most tissues, whereas DS-epi2 is the main epimerase in the brain [21,22]. DS-epi2 also has a relatively high expression in the kidney.

The epimerase reaction is reversible, with an equilibrium of 9 : 1 (GlcA to IdoA) under in vitro conditions when the biosynthetic complex has been solubilized with detergent [4]. On the other hand, CS/DS chains in vivo can contain a higher proportion of IdoA. This is assumed to be achieved through functional collaboration between DS-epi1 and D4ST1 (Fig. 3) [23]. In support of this, transient down-regulation of D4ST1 results in a reduced IdoA content [24]. Genetic mutations in D4ST1 found in a new type of Ehlers–Danlos syndrome (i.e. adducted thumb-clubfoot syndrome) also result in CS/DS of low IdoA content [25].

Little is known about the regulation of epimerase activity. Transforming growth factor (TGF)-β-stimulated fibroblasts have reduced levels of epimerase activity, a reduced expression of D4ST1 and an increased expression of C4ST1, resulting in CS/DS with a considerably lower amount of IdoA [26]. This effect is further increased by combined treatment with TGF-β, epidermal growth factor and platelet-derived growth factor (PDGF) (9). In another study, PDGF promoted the migration of fibroblasts, comprising a mechanism that is proposed to involve the up-regulation of IdoA in the proteoglycan CD44 [27].

The products of DS-epi1 and 2 are difficult to assess as a result of the complex interaction with D4ST1. DS-epi1 can generate long blocks of IdoA together with D4ST1 (Fig. 3). Down-regulation of D4ST1 resulted in the abrogation of IdoA-containing blocks without affecting overall epimerase activity [24]. The role of DS-epi2 has been more difficult to assess. Overexpression of DS-epi2 increased IdoA in hybrid structures (Fig. 3). No increase of IdoA blocks was recorded upon overexpression of DS-epi2, whereas overexpression of DS-epi1 resulted in enhanced block formation [16]. By contrast, down-regulation of DS-epi2 in fibroblasts decreased the proportion of IdoA blocks, although to a smaller degree than that obtained by down-regulation of DS-epi1. Data obtained from DS-epi1 knockout mice show that DS-epi2 mainly forms alternating structures [28]. These data indicate that DS-epi2 might be primarily involved in the formation of isolated or alternating IdoA structures (Fig. 3).

Different proteoglycans produced by the same cell can vary greatly with respect to their IdoA content and distribution. For example, decorin and biglycan have been found to contain blocks of IdoA, whereas versican only has isolated IdoA. Other studies have suggested that the core protein regulates the activity of the DS epimerases. This was demonstrated by the generation of chimeric proteins of decorin, which has a high content of IdoA, and colony-stimulating factor, a part-time proteoglycan with a low content of IdoA. The chimeric decorin–colony-stimulating factor contained less IdoA than the unmodified decorin [29]. This suggests that core proteins carry information that may direct the proteoglycan cores to compartments within the Golgi complex with different amounts of DS epimerase activity [30].

**Functions of IdoA as indicated by targeting of the two epimerases**

The phenotype observed in DS-epi1 knockout mice is dependent upon the genetic background. Using mice with a pure C57BL6 genetic background, all pups die perinatally, whereas, when using mice with a pure NFR background, approximately half of the pups die. The NFR pups have a retarded growth rate in the late
embryological stages of development and, furthermore, ~20% of the pups display gastroschisis, an abdominal wall-closure defect that presents intestines outside the body (R. Gustafsson, unpublished data). DS-epi1 depleted mice in a mixed 129Sv/C57BL6 genetic background have been investigated in more detail. The pups were born at a normal Mendelian frequency [28]. At birth, they are smaller and have a crooked tail. Because decorin is a major proteoglycan involved in the organization of collagen fibrils in skin, this tissue was studied in more detail. DS-epi1−/− skin was more fragile than the skin of wild-type mice. DS-epi1−/− collagen fibrils were more heterogeneous in denaturation profiles and in vitro experiments showed that, in DS-epi1−/− skin, decorin was the proteoglycan that was responsible for altered collagen structure (Fig. 4A). Electron microscopy showed that the diameter of DS-epi1−/− fibrils was 85 nm compared to 62 nm for wild-type mice [28]. In summary, iduronic acid in the CS/DS chain and particularly of IdoA blocks participates in skin collagen fibril maturation.

DS-epi2−/− mice do not show any evident phenotype [22]. The brain was analyzed because DS-epi2 is the predominant epimerase in this tissue [22,31]. Accordingly, DS-epi2−/− brains had a 90% reduction in epimerase activity. The brains of newborn mice contain little IdoA (2% of the total chain), and this was further reduced in DS-epi2−/− mice. However, the brain extracellular matrix (ECM) architecture was unaltered. It would be interesting to determine whether more subtle phenotypes such as behavioural changes are present in DS-epi2−/− mice.

Mice deficient in DS-epi1 and 2 were recently obtained in a mixed 129Sv/C57BL6 genetic background. A large proportion of the pups die perinatally, although a few survive until 7 weeks of age. Double ERK1/2

Leukocyte

NF-κB

ICAM-1

Fig. 4. Overview of the functions of IdoA in CS/DS. Role of IdoA in the storage of cytokines growth factors and collagen fibril formation (A), Borrelia infection (B), atherosclerosis (C), coagulation (D), P-selectin-dependent leukocyte recruitment (E), activation of cytokine and growth factor receptors (F) and leukocyte recruitment by ICAM (G).
knockout mice are dwarf and have approximately half the size and weight of their wild-type littermates.

Down-regulation of DS-epi1 has been achieved in the frog *Xenopus laevis* using morpholino injections (E. Pera, unpublished data). Several abnormalities were observed, such as the absence of the dorsal fin, which could be explained by the altered migration of neural crest cells into that anatomical structure.

**Genetic alterations affecting IdoA formation in humans**

There are no mutations in DS-epi1 associated with human diseases. However, mutations in D4ST1, which functionally collaborates with DS-epi1 to make IdoA blocks (Fig. 3), result in adducted thumb-clubfoot syndrome [32]. Mutations of D4ST1 result in reduced amount of IdoA in CS/DS [25], also resulting in a defect in collagen fibril maturation and reduced collagen strength [28]. This autosomal recessive syndrome [33] is characterized by facial changes, contractures of thumbs and fingers, joint instability, skin hyperextensibility, and heart and kidney defects. Additionally, myopathy has been described in these patients [34].

DS-epi2 has been genetically associated with bipolar disorder, which is a disease affecting ~1% of mankind [15]. Interestingly, two single nucleotide polymorphisms predicted to change the amino acid sequence were present in the bipolar disorder group and not in the control group.

**The role of IdoA in stem cell development**

Embryonic stem cells are obtained from embryos and can be maintained in cell cultures as pluripotent stem cell lines with a capacity to differentiate into whole embryos. Studies have shown a four- to six-fold increase of CS/DS during the differentiation of murine embryonic stem cells to embryoid bodies and to extra embryonic endodermal cells. The formation of embryoid bodies and extra embryonic endodermal cells was accompanied by a two- and four-fold increase of IdoA, respectively [35], suggesting a role for IdoA. The biosynthetic genes *DSE, DSEL* and *CHST14*, coding for D4ST1, were expressed at all stages. *CHST14* was also expressed in the extra embryonic endodermal cells. However, the detailed structure of CS/DS, as well as its functions, still needs to be determined.

CS/DS is enriched in the neural stem cell niche and has been shown to play important role in the differentiation of neural progenitor cells [36]. Its importance has been demonstrated in progenitor cells from mice with ablations of C4ST1 (a 4-O-sulfotransferase acting on GlcA-containing sequences) and D4ST1. Down-regulation of D4ST1 resulted in the abrogation of IdoA blocks, as well as decreased neurogenesis and proliferation and a change in the expression of cell surface receptors for fibroblast growth factor (FGF)-2 and epidermal growth factor, whereas C4ST1 deficiency did not affect these processes [37]. The importance of IdoA motifs was further underlined by the fact that mRNA expression of the DS epimerases was higher in differentiated neurones than in precursor stem cells [38].

**IdoA-containing structures in brain development**

CS/DS structures are implicated in brain development [39] and injury to the central nervous system [40]. During development, IdoA-containing structures (iA, iB, iE and iD) are ubiquitous in different parts of the brain [31,41], although at low concentrations. Indeed, CS/DS brains of newborn mice comprise only 2% iduronic acid [22]. The CS/DS bioenzymatic machinery is carefully regulated during brain development, resulting in a large variation of IdoA-containing structures. For example, in the cerebellum, iD decreases and iB increases from newborn to adult age [31]. Interestingly, the embryo-derived CS/DS shows a greater binding of FGFs (FGF-2, -10 and -18), pleiotrophin, midkine, vascular endothelial growth factor (VEGF) and hepatocyte growth factor (HGF) than CS/DS from the brains of adult animals [42].

**The role of IdoA in CS/DS under pathological conditions**

**Inflammation**

The involvement of CS/DS in inflammation has been extensively explored, whereas the role of IdoA is not well defined [43,44]. The inflammatory response initiated by infection or injury results in diverse processes, involving cell recruitment, extravasation and cell/pathogen clearance. For example, during wound healing, CS/DS is reported to be the dominating GAG in wound fluid [45,46]. FGF2 and FGF7 are two important growth factors during wound repair and they have been shown to preferentially bind to IdoA-containing motifs in CS/DS, promoting proliferative processes (Fig. 4A,F). CS/DS has been proposed, in combination with FGF-10, as a pharmacological accelerator of wound closure as a result of its capacity to stimulate re-epithelialization [47]. CS/DS can potentially affect several steps during cell recruitment.
For example, CS/DS has been shown to interact with P-selectin, which is expressed on endothelial cells and platelets [48] (Fig. 4E). CS/DS is reported to influence the recruitment of polymorphonuclear cells in a thiglycollate-induced inflammatory model in a supposedly P-selectin manner [49]. RANTES, a leukocyte-recruiting chemokine, also interacts with IdoA-containing segments in CS/DS [50]. An essential step during extravasation is the increased expression of intercellular adhesion molecule-1 (ICAM-1) on endothelial cells. IdoA in CS/DS induces endothelial expression of ICAM-1 mediated by nuclear factor-kB [45] (Fig. 4G). Interestingly, macrophages are reported to produce CS/DS containing up to 70% of IdoA [51]. After lipopolysaccharide stimulation, macrophages predominantly secrete CS/DS, either as free chains or bound to the serglycin core protein.

**Immune response**

Autoimmunity is a result of a disarray in the immune response, which becomes directed towards its own tissue and cells. B cells participate in autoimmunity by the production of antibodies and presentation of self-antigens to T cells. IdoA motifs in CS/DS are reported to augment the proliferation of B1-a cells and increase their autoantibody production [52]. IdoA in CS/DS interacts with components from apoptotic and dead cells and forms complexes that enhance autoantibody production. IdoA-containing structures in CS/DS bind autoantigens, which were enriched after CS/DS-affinity chromatography of cellular lysates. Two hundred autoantigens were identified by MS and could be used in western blot experiments to detect different autoantibody patterns of diagnostic value in patient sera [52,53]. Further studies are needed to clarify the physiological role of CS/DS in the generation of natural autoantibodies.

**Atherosclerosis**

Atherosclerosis, an inflammatory-driven disease, is characterized by the accumulation of cholesterol in arterial blood vessels, resulting in thicker and more fragile artery vessels. Binding of low-density lipoprotein (LDL) to GAGs is considered to be one of the steps in the onset of this disease [54]. The GAG interaction enhances LDL uptake by macrophages, leading to the formation of foam cells (Fig. 4C). IdoA both in CS/DS and heparan sulfate is reported to enhance the binding of VLDL and LDL [55,56]. Recently, it was reported that an antibody against CS/DS inhibited the LDL–CS/DS interaction and inhibited LDL oxidation in vitro [57]. Furthermore, the injection of anti-CS/DS antibody in an atherosclerosis model of ApoE−/− mice resulted in decreased arteriosclerotic lesions [58].

**Coagulation**

Coagulation is essential under normal physiological conditions and several pathological conditions (e.g. cancer, atherosclerosis and sepsis) have enhanced coagulation. Thrombin, a serine protease, catalyzes the conversion of fibrinogen to fibrin, which forms blood clots in conjunction with platelets. Heparin cofactor II (HCII) is a thrombin inhibitor and the only known serpin to be activated by IdoA-containing CS/DS (Fig. 4D). The HCII binding site to CS/DS differs from that to HS [59]. The HCII binding structures in CS/DS contain IdoA-2S-GalNAc-4S [60] or GlcA-GalNAc-4,6-disulfated [61] in hexa- and octasaccharides as minimal binding motifs. The complex CS/DS-HCII is considered to be the major anticoagulant system after injury of the vessel wall [60,62,63]. CS/DS containing 2-O-sulfated IdoA also controls coagulation by activating protein C [64].

**Infection**

CS/DS is involved in bacterial infections. *Borreilia* (causing Lyme disease) was shown to use the core protein of decorin, as well as its CS/DS side chain, as a binding target in the initial phase of infection [65] (Fig. 4B). CS/DS released from decorin by proteases produced by *Pseudomonas, Enterococcus* and *Streptococcus* [66] targets α-defensin and inhibits its bactericidal activity. The optimal structure for interaction to α-defensin is a motif containing a mix of IdoA and GlcA, which is found in decorin present in fibrous connective tissue [66].

**IdoA motifs in cancer**

CS/DS is implicated in several cancer-promoting processes, such as cell proliferation and metastasis [3]. DS-epi1, previously named SART2 (squamous cell carcinoma antigen recognized by T cell 2), is highly expressed in many tumours and cell lines [20]. DS-epi1 expressed by cancer cells was recognized by HLA-A24-restricted and tumour-specific cytotoxic lymphocytes. Peptides from DS-epi1 were used in peptide-based immunotherapy phase I clinical trials for prostate cancer [67], glioblastoma multiforme [68] and hepatocellular carcinoma [69] with moderate success. We have established that DS-epi1 is not tumour specific because DS-epi1 is ubiquitously expressed in normal tissues [21]. Squamous cell carcinoma from oesophagus
contains epimerase activity that is increased four- to five-fold compared to normal oesophagus [13]. DS-epi1 is localized both in stroma surrounding the tumour and in cancer cells. To investigate the role of IdoA, DS-epi1 was stably down-regulated in oesophage squamous carcinoma cell lines using shRNA sequences. IdoA was shown to facilitate the binding of HGF to its receptor and was essential for cMET-dependent signalling [13] (Fig. 4F). In addition, DS-epi1 down-regulated cells displayed fewer cytoplasmic stress fibres than control cells. Furthermore, the focal adhesion complexes were evenly distributed at the cell surface in DS-epi1 down-regulated cells compared to control cells, which displayed focal adhesion complexes predominantly at the leading edge. This resulted in less migration and invasion of DS-epi1 down-regulated cells compared to control cells [13].

Different CS/DS structures mediate diverse function during cancer development. The sulfation pattern of CS/DS in cancer differs from normal tissue. For example, 6-O-mono-sulfated disaccharides are accumulated in tumours compared to normal tissues, whereas 4-O-mono-sulfated disaccharides are reduced [70]. During metastasis, CS/DS disaccharides sulfated at positions 4 and 6 (E units) present on the surface of cancer cells facilitate colonization of the lung and liver [71,72]. The process might be mediated by the receptor RAGE, which is highly expressed in the lung [73]. Another pro-metastatic activity of the E units on cancer cells could be a result of the capability to bind platelet P-selectin [49], resulting in the formation of tumour microemboli. These cell–platelet aggregates protect cancer cells against elimination by the immune system. IdoA in CS/DS is also essential to mediate selectin binding. Two CS/DS structures containing IdoA (iB units or iD units), as isolated from marine animals, inhibit metastasis in a P-selectin-dependent manner in a metastatic tumour model [49]. Several studies report that CS/DS structures mediate growth factor and chemokine binding. IdoA is essential for HGF-mediated binding and an IdoA-containing tetrasaccharide is the minimum structure required to confer affinity [74]. Exogenously added IdoA-containing motifs inhibit the proliferation of normal and malignant cells [75]. Elimination of CS/DS on the cancer cell membrane by chondroitinase B inhibits the migration and invasion of tumour cells [76].

**Future perspectives in research and clinical therapy**

Still largely unknown is how the complex structure of CS/DS is formed and how it is regulated. A key question is the organization of the biosynthetic enzymes in the Golgi and how this organization is modulated in different cells and tissues. The role of the two different epimerases, DS-epi1 and 2, as well as that of D4ST1, needs to be clarified.

Different functions of IdoA have been found both in vitro and in vivo. The human situations where DS-epi1 expression is changed in tumours and where D4ST1 mutations lead to deranged connective tissue have been highlighted. The importance of IdoA is evident from observations of DS-epi1 KO mice, which die perinatally and/or present gastrointestinal. Furthermore, a decrease of IdoA leads to an altered collagen structure, resulting in a decreased tensile strength. Provocation of mice with targeted DS-epi1 and 2 will most likely provide more information about other biological functions of IdoA. Other data indicate the importance of IdoA in cytokine activity and storage, cell proliferation and migration, the control of coagulation, the formation of autoantibodies, the control of stem cell stability and differentiation.

In disease, IdoA contributes to cancer progression and infection. New avenues for future therapies have been tested, such as vaccination against cancer [67–69], or are warranted to control infection [65,66] and cancer [13,76,77]. DS epimerases inhibitors could be used in cancer and fibrosis, as well as to guide stem cell differentiation [3].

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