The structure of the Shiga toxin 2a A-subunit dictates the interactions of the toxin with blood components

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Abstract
Hemolytic uremic syndrome (eHUS) is a severe complication of human infections with Shiga toxins (Stxs)–producing Escherichia coli. A key step in the pathogenesis of eHUS is the interaction of Stxs with blood components before the targeting of renal endothelial cells. Here, we show that a single proteolytic cleavage in the Stx2a A-subunit, resulting into two fragments (A1 and A2) linked by a disulfide bridge (cleaved Stx2a), dictates different binding abilities. Uncleaved Stx2a was confirmed to bind to human neutrophils and to trigger leukocyte/platelet aggregate formation, whereas cleaved Stx2a was ineffective. Conversely, binding of complement factor H was confirmed for cleaved Stx2a and not for uncleaved Stx2a. It is worth noting that uncleaved and cleaved Stx2a showed no differences in cytotoxicity for Vero cells or Raji cells, structural conformation, and contaminating endotoxin. These results have been obtained by comparing two Stx2a batches, purified in different laboratories by using different protocols, termed Stx2a(cl; cleaved toxin, Innsbruck) and Stx2a(uncl; uncleaved toxin, Bologna). Stx2a(uncl) behaved as Stx2a(cl) after mild trypsin treatment. In this light, previous controversial results obtained with purified Stx2a has to be critically re-evaluated; furthermore, characterisation of the structure of circulating Stx2a is mandatory to understand eHUS–pathogenesis and to develop therapeutic approaches.

1 | INTRODUCTION

Infections with enterohemorrhagic Escherichia coli (EHEC) are the major cause of EHEC-associated hemolytic uremic syndrome (eHUS), the most common reason for acute renal failure in childhood with a mortality rate of 3–5% (Rosales et al., 2012; Tarr, Gordon, & Chandler, 2005). However, only 5–15% of patients infected with EHEC develop eHUS with the characteristic symptom triad of microangiopathic hemolytic anaemia, thrombocytopenia, and acute renal failure (Tarr et al., 2005).

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Shiga toxins (Stxs) represent major EHEC virulence factors in eHUS (Karch, Tarr, & Bielaszewska, 2005). Two different types of toxin, namely, Stx1 and Stx2, are known. Within each of the toxin types, several subtypes have been identified and subtype 2a was shown to correlate significantly more with severe illness in humans, such as eHUS (Friedrich et al., 2002; Orth et al., 2007). Stxs, which belong to the family of ribosome inactivating proteins, are ABS holotoxins that consist of one enzymatically active A moiety (~32 kDa) non-covalently associated with a pentameric B-subunit (~7.7 kDa per monomer; Paton & Paton, 1998). The pentameric B-subunit is responsible for binding to the globotriaosyl-ceramide (Gb3Cer) or Gb4Cer cellular receptors (Bauwens et al., 2013), mainly expressed by human renal cells such as glomerular endothelial cells (Obrig, 2010).

After internalisation, Stx is transported from early/recycling endosomes to the Golgi apparatus (Garred, van Deurs, & Sandvig, 1995; Mallard et al., 1998) and then to the endoplasmatic reticulum (ER; Sandvig et al., 1992). During the transport, the A-subunit is cleaved by the protease furin resulting in two fragments, A1 and A2 (~27.5 kDa and ~4.5 kDa, respectively). The A1-fragment remains linked to the A2-fragment by a disulfide bond between C241 and C260 (Garred et al., 1995) and A2 interacts with the pentameric ring formed by the B subunits. The loop between the two cysteines contains the sequence [Arg·X·X·Arg], the consensus motif for cleavage by furin, which is also recognised by trypsin (Garred et al., 1995). In the ER, the A1 subunit is released from the A2-B5 complex by reduction of the disulfide bond. The A1 subunit is translocated from the ER into the cytosol or into the nucleus where it inhibits protein synthesis and induces the formation of apurinic sites in chromatin (Bergan, Dyve Lingelem, Sinn, Skotland, & Sandvig, 2012; Brigotti et al., 2002; Sandvig & van Deurs, 2005; A. Suzuki et al., 2000).

Before targeting cells in the kidney, Stx binds to blood components, but it is not clear whether the A subunit is still intact then or already cleaved by bacterial or host proteases, and such interactions are involved in the pathogenesis of eHUS. In particular, Stxs are binding to blood cells, like neutrophils, monocytes, or platelets (Brigotti et al., 2008; Brigotti et al., 2013; Cooling, Walker, Gille, & Koerner, 1998; Karpman et al., 2001), and subsequent activation has been reported. As a consequence, formation of leukocyte-platelet aggregates occurs, followed by the release of 1-μm-microvesicles bearing Stxs and other factors such as tissue factor and activated complement factors (Stahl, Sartz, & Karpman, 2011; Stahl, Sartz, Nielsson, Bekassy, & Karpman, 2009). Furthermore, Stx has been demonstrated to activate complement in vitro and to bind factor H (FH; Orth et al., 2009).

Here, we show that, depending on the presence or absence of a single proteolytic cleavage in the disulfide bridged loop of the A subunit, Stx2a has different binding properties for key circulating components involved in the pathogenesis of HUS (neutrophils, FH) and different biological properties (formation of leukocyte-platelet aggregates). These results have been obtained by comparing two different Stx2a batches purified in two laboratories by using different protocols. This is in line with the previously reported controversial results regarding the effects of purified Stxs.

2 | RESULTS

2.1 | Characterisation of Stx2a

Because preliminary data from the Innsbruck laboratory did not confirm Stx2a binding to human neutrophils as previously demonstrated in the Bologna laboratory (Brigotti et al., 2013), we performed a comparative analysis of the biological activity and of the physicochemical properties of the different toxin preparations purified in Innsbruck (Austria) and in Bologna (Italy).

Sequencing analysis confirmed that for both preparations, the stx2a genes were identical for the A subunits (Figure S1). The B subunits showed one mismatch at position 36 (Figure S1), which, however, did not change the protein sequence.

The amount of lipopolysaccharide (LPS) in the Stx2a preparations (45 Eu/ml, 56 Eu/mg, Innsbruck; 35 Eu/ml, 67 EU/mg of protein, Bologna) measured by the Limulus assay was comparable.

SDS-PAGE analysis performed at non-reducing conditions revealed the presence of two Coomassie-stained bands corresponding to A and B subunits in Stx2a prepared in Innsbruck (Figure 1a), whereas an additional band appeared at reducing conditions (Figure 1a) having the apparent molecular mass of the A1 fragment (27.6 kDa) and migrating faster than the A subunit (31.7 kDa).

Densitometric analysis showed that Stx2a purified in Innsbruck consists of 16% A chain and of 84% A1 fragment. Because the A1 fragment was observed only in the presence of mercaptoethanol as reducing agent (Figure 1a), we conclude that the A subunit is cleaved in the loop between C241 and C260 and that the two resulting fragments A1 and A2 are linked by an intact disulfide bond (Figure 1b). The cleaved Stx2a from Innsbruck was named Stx2a(cl) throughout the paper. Conversely, Stx2a prepared in Bologna and run at reducing and non-reducing conditions (Figure 1a) showed a single 31.7 kDa band for the intact A subunit and, hence, was named Stx2a(uncl). Treatment of Stx2a(uncl) with furin, the cell protease involved in the intracellular activation of Stxs (Garred et al., 1995), or with trypsin (Donohue-Rolfe, Jacewicz, & Keusch, 1988), which mimics the single proteolytic cleavage of the Stx A subunit occurring in cells, reproduced the pattern observed in Stx2a(cl) obtaining the A1-fragment (27 kDa; Figure 1c).

2.2 | Cleaved and uncleaved Stx2a differ in the ability to bind human neutrophils

Stx2a(uncl) was found to bind human neutrophils by indirect flow cytometric analysis (Figure 2), giving full saturation of receptors at 60 nM concentration (Figure 2a), whereas Stx2a(cl) and trypsin-cleaved Stx2a were not recognised by human neutrophils (Figure 2b,d). This was confirmed with neutrophils from three different healthy donors (Figure 2b). Consistently, furin treatment of Stx2a impaired neutrophil binding to extents (60.8 ± 8.6% inhibition with respect to untreated Stx2a(uncl), n = 3, P = 0.0066) related to the amount of generated A1 fragment (47%).
To achieve a more physiological situation, binding of Stx2a to neutrophils was also investigated in human blood. Binding of cleaved or uncleaved toxins to neutrophils was not influenced by the presence of blood components (Figure 2c).

### 2.3 Cleaved and uncleaved Stx2a differ in the ability to form leukocyte/platelet aggregates in human blood

Although Stx2a(uncl) was fully active with respect to leukocyte/platelet aggregates formation, Stx2a(cl) and trypsin-cleaved Stx2a did induce lower percentages of leukocyte/platelet aggregates both in representative experiments (Figure 3a,b) and in confirmation with three healthy donors (Figure 3c,d). Addition of LPS amounts present in Stx2a preparations to blood produced no aggregates, as well as the addition of heat-treated Stx2a(uncl).

### 2.4 Cleaved and uncleaved Stx2a differ in the ability to interact with complement factor H

Binding of Stx2a to complement factor H, assessed by ELISA, was confirmed for Stx2a(cl) and was also detected for trypsin-cleaved Stx2a but not for Stx2a(uncl; \( P < 0.001 \) for all; Figure 4). FH binding was also evaluated with LPS (amount present in Stx2a preparations) and was found to have no effect (data not shown). Thus, the binding patterns of the different toxin forms to FH was opposite (cleaved Stx2a active, uncleaved Stx2a inactive) with respect to aggregate formation or interaction with granulocytes (cleaved Stx2a inactive, uncleaved Stx2a active).

### 2.5 Cleaved and uncleaved Stx2a do not activate human platelets

It has been previously reported that Stx can activate platelets (Karpman et al., 2001), hence, concurring to the development of thrombocytopenia during eHUS. Both Stx2a(uncl) and Stx2a(cl) were unable to activate platelets, which conversely are sensitive to thrombin used as positive control (Figure S2) as assessed by increased surface expression of CD62P.

### 2.6 Cleaved and uncleaved Stx2a are fully active toxins for Gb3Cer-expressing cells

Although differing in neutrophil binding, aggregate formation and FH-binding abilities, the different toxin forms were found to be fully active and similar in inhibiting translation in sensitive cells endowed with the receptor Gb3Cer such as Raji cells (Figure 5a). After a short incubation with Stx2a, translation was determined in the presence of a radioactive amino acid as previously described (Arfilli et al., 2015). The calculated IC\(_{50}\) were very similar and as expected for fully active Stxs (Brigotti et al., 2002; Brigotti et al., 2007; Brigotti et al., 2011). It has been shown that the presence of human serum strongly reduced the inhibitory power of Stx2a on this system because of the presence of HuSAP (Arfilli et al., 2015). This circulating protein complex, by capturing Stx2a, prevents the targeting of Gb3Cer-containing cells by the toxin (Armstrong et al., 2006; Kimura, Tani, Matsumoto Yi, & Takeda, 2001; Marcato, Vander Helm, Mulvey, & Armstrong, 2003). When adding human serum, translation inhibition was similarly strongly impaired for cleaved or uncleaved toxins (Figure 5b). The cytotoxicity
profiles on Vero cells, chosen to reveal long-term cytopathic effects, were also found to be similar (Figure 6).

2.7 Cleaved and uncleaved Stx2a do not differ in conformation

To gain information on the effect of cleavage on the conformation of the toxins, Stx2a(cl) and Stx2a(uncl) were examined by circular dichroism (CD) and fluorescence. The CD spectra of the two proteins were almost identical indicating conformational similarity (Figure 7a). The spectrum exhibits a negative minimum at 220 nm and a less intense slightly positive band above 235 nm. According to the CD spectral profiles of proteins of known structure (Brahms & Brahms, 1980), we can conclude that both contain a significant fraction of $\alpha$-helix structure and only a small fraction of $\beta$-sheet structures.
This was further confirmed by their fluorescence behaviour (Figure 7b). Stx2 possesses 12 tryptophan (Trp) residues, the dominant fluorophores in proteins, two of them in the A subunit and two in each of the five B subunits. The fluorescence spectra of the two proteins were very similar in shape and peak at 347 nm, a quite red shifted value indicative of an average polar environment for Trp residues in both proteins. The maximum in the excitation spectra at 281, as well as the shape, corresponds to Trp absorption in polar environment. Global analysis of the decay profiles for each protein evidenced two lifetime ($\tau_1$) components that were almost the same for the two proteins (Stx2a(d)): $\tau_1 = 2.64$ ns, $\tau_2 = 7.22$ ns; Stx2a(uncl)): $\tau_1 = 2.50$ ns, $\tau_2 = 7.33$ ns) and have similar fractional contributions. The presence of two lifetime components, similar in both protein samples, points to the presence of at least two types of Trp environments of different average polarity.

The photophysical study allowed to conclude that no significant conformational changes occurred in the two proteins in spite of the differences observed in their structure.

3 | DISCUSSION

Stxs are believed to be the main virulence factors in eHUS, and amongst the Stx subtypes, Stx2a is strongly associated with severe disease, including eHUS (Orth et al., 2007). Thus, several in vitro studies have been performed with purified Stx2a to elucidate the pathomechanism of this most extreme outcome of EHEC infection. However, for Stx purification, there is no standardised protocol available and, thus, different and often poorly detailed Stx purification methods, which employ different Stx-producing strains have been used.

FIGURE 3  Stx2a(cl) or trypsin‐cleaved Stx2a added to human blood are not efficient in triggering the formation of leukocyte/platelet aggregates. Whole human blood was treated with 1 nM Stx2a(cl), Stx2a(uncl), or trypsin‐cleaved Stx2a. When indicated lipopolysaccharide amount (0.01 EU) similar to that contaminating Stx2a preparations was added in the assay. Then, the formation of neutrophil/platelet or monocyte/platelet aggregates was assessed by direct flow cytometric analysis. Values obtained in a representative experiment expressed as percentage of aggregates on the neutrophil (a) or monocyte (b) populations; data are means ± SD (n = 2). (c, d) Values minus the appropriate controls (no treatment or trypsin + trypsin inhibitor) from three independent experiments were used to calculate the percentage of total aggregates with respect to Stx2a(uncl); data are means ± SD (n = 3). After incubation of human blood with Stx2a(uncl), the percentages of neutrophil/platelet aggregates and of monocyte/platelet aggregates on the whole population of neutrophils or monocytes were 55.0 ± 8.0% and 47.5 ± 14.2% (mean ± S.D., n = 3), respectively. *P < 0.05, **P < 0.01, ****P < 0.0001 with respect to Stx2a(uncl) are indicated in the figure.
Different capabilities of Stx2a have been described that might contribute to the pathogenesis of eHUS. These characteristics include cytotoxicity against various cells, often using Vero cells as model, protein synthesis inhibition, neutrophil binding, platelet binding and activation, formation of leukocyte/platelet aggregates, binding to different serum proteins, such as HuSAP or complement factor H, and complement activation (Bauwens et al., 2011; Brigotti et al., 2013; Brigotti et al., 2008; Cooling et al., 1998; Karpman et al., 2001; Lentz, Leyva-Illades, Lee, Cherla, & Tesh, 2011; Marcato et al., 2003; Orth et al., 2009; K. Poolpol et al., 2014). However, several of these Stx2a-induced features have been controversially discussed. Either the Stx2a-induced effect was not comparable in magnitude, or some of these effects could not even be reproduced by other groups.

Investigation of Stx2a-induced capabilities of Stx2a(uncl) or of cleaved forms (Stx2a(cl) and trypsin-cleaved Stx2a) revealed comparable results regarding cytotoxic activity and absent activation of platelets for both preparations. Conversely, in the case of key Stx2a-induced capabilities, striking differences were found: Stx2a(uncl) was confirmed to bind to human neutrophils and to trigger leukocyte/platelet aggregate formation whereas cleaved Stx2a, such as Stx2a(cl) and trypsin-cleaved Stx2a, was ineffective. In contrast, binding to FH was confirmed for Stx2a(cl) and was also detected for trypsin-cleaved Stx2a; Stx2a(uncl) failed to bind FH.

Limulus assay revealed no significant differences in the low-LPS amount that contaminated Stx preparations. Moreover, running experiments (neutrophil binding, leukocyte-platelet aggregate formation, and FH binding) with these LPS amount did not show any effect.

Structure analyses of the Stx2a preparations revealed conformational similarity of both toxins including a comparable content of α-helices and β-sheet structures. However, SDS-PAGE analyses showed that Stx2a(cl) A subunit is cleaved in two fragments whose molecular

**FIGURE 4**  Stx2a(cl) and trypsin-cleaved Stx2a binds to complement FH. Immobilised Stx2a(cl) or Stx2a(uncl) or trypsin-cleaved Stx2a (2 μg/well each) was incubated with FH (2 μg). As controls trypsin + trypsin inhibitor (PMSF) or BSA were incubated with FH. Additional controls were performed by incubating the proteins with buffer (TBST) instead of FH. As positive control FH was used. The binding capacity of Stx2a(cl) and trypsin-cleaved Stx2a to FH were significantly higher compared with Stx2a(uncl) and all negative controls (P < 0.001 for all). ***P < 0.001, ****P < 0.0001

**FIGURE 5**  Inhibition of protein synthesis in Raji cells by Stx2a(cl), Stx2a(uncl), and trypsin-cleaved Stx2a. Raji cells were treated 3 hr with different concentrations of toxins in the absence (a) or in the presence (b) of 10% human serum then protein synthesis was measured in the presence of labelled leucine. IC50 was calculated by the linear regression between fractional activity and the log of Stx2a concentrations

**FIGURE 6**  Cytotoxicity profiles of Stx2a(cl) and Stx2a(uncl) on Vero cells are very similar. Vero cells were incubated 48 hr with a broad range (10 fg/ml-10 μg/ml) of Stx2a concentrations (150 aM-150 nM). Residual cell density was measured by crystal violet method, and the percentage of living cells is shown on the y-axis.
masses correspond to A1 and A2 fragments. The A fragments of Stx2a(cl) are linked by a disulfide bond, shown by the fact that only the mercaptoethanol-reduced gels resolved A1 and A2 fragments, whereas use of non-reducing sample buffer revealed only one band for the A subunit. Cleavage of the A subunit was found to happen early in the purification procedure of Stx2a(cl) at the sonication step (data not shown); this is in agreement with the fact that this step is lacking in the purification method yielding the Stx2a(uncl). The structural change was reproduced by treating Stx2a(uncl) with trypsin and furin to mimic the cleavage observed during the Stx2a(cl) purification.

Stx2a has been described to bind to human neutrophils via its A subunit at TLR4 (Brigotti et al., 2013). The fact that Stx2a(cl) and trypsin-cleaved Stx2a did not interact with human granulocytes suggests that the binding site for neutrophils might be located close to the cleaved area of the A subunit but not in the active site of Stx2a, as cytotoxic activity and protein synthesis were unchanged. Furthermore, the fact that Stx2a(uncl) induced a significantly higher percentage of neutrophil and monocyte-platelet aggregate formation than Stx2a(cl) might indicate an important role of TLR4 in these phenomena. It should be noted that all the actors involved (platelets, monocytes, and neutrophils) express this receptor. The Stx2a B subunit has previously been suggested to contain the binding site for FH (Poolpol, 2012). We hypothesize that nicking of the A subunit might influence the quaternary structures of the whole toxin affecting A/B-subunit interactions, thus explaining the different abilities of Stx2a preparations concerning the binding to FH. However, it seems worth to note that cytotoxic activity of both toxin preparations was comparable indicating that the receptor binding sites located in the B subunit (Bauwens et al., 2013) of Stx2a were not affected and that also the cooperation between B and A chains (binding to Gb3Cer and intracellular enzymatic action, respectively) was operative for both toxin batches.

Cleavage of the Stx2a A subunit has been described during the cellular transport of Stx2a; the A1 fragment is released in the ER and subsequently conveys protein synthesis inhibition at the ribosome (Sandvig et al., 1992). However, nicking of the A subunit might also occur in the intestinal mucus (Melton-Celsa, Darnell, & O’Brien, 1996) or is likely induced by bacterial proteases released by lysed EHEC during infection. In this case, one can assume the presence of different proportion of cleaved and uncleaved Stx2a in blood during the natural course of EHEC infections and in the transition from hemorrhagic colitis to eHUS. There are very limited data showing presence of Stx2a in blood of EHEC-infected patients. Lopez and co-workers have detected free Stx2a in sera of three EHEC-infected patients during the prodromal intestinal phase before the onset of eHUS (Lopez et al., 2012). He and co-author have found only very low amounts of Stx2a in sera of eight patients with overt eHUS (He et al., 2015). Arfilli and co-workers have demonstrated presence of functional active Stx2a in sera of three EHEC-infected patients during hemorrhagic colitis (Arfilli et al., 2015). However, none of these studies presented any data on the structure and/or conformation of Stxs in the blood of eHUS patients. Thus, it is not clear at which step of eHUS pathogenesis Stx2a cleavage could occur and to which extent.

Furthermore, it cannot be stated whether and to what extent neutrophil binding, leukocyte-platelet aggregation, and factor H binding occur and contribute to eHUS pathogenesis. Interestingly, uncleaved and cleaved Stx2a seem to participate similarly in HuSAP-related effects as supported by our experiments in the presence of human blood and serum. This blood protein prevents the targeting of Gb3Cer-expressing cells by interacting with Stx2a and, at the same time, promotes the interaction of the toxin with human neutrophils and subsequent steps involved in the pathogenesis of eHUS.

We actually speculate that for the binding to neutrophils, it is mandatory that the toxin remains uncleaved because these cells do not express Gb3Cer. Then, during the subsequent pathogenetic steps (formation of aggregates and release of microvesicles) or in the vicinity of the target organ, the toxin could get cleaved, allowing the interaction with factor H. Moreover, circulating cells co-expressing TLR4 and Gb3Cer, such as monocytes and platelets, could interact with cleaved (through Gb3Cer) or uncleaved (through TLR4 and Gb3Cer) Stx2a. Hence, the relative proportion of cleaved and uncleaved Stx2a on these cells could determine (a) different amounts of the deriving aggregates and microvesicles and (b) different pathogenic factor contents (activated complement components) in aggregates and
microvesicles. Finally, cleavage of cell surface-associated Stx2a could represent a bound/unbound mechanism allowing toxin delivery to target endothelia after transport by TLR4-expressing circulating cells.

In conclusion, this study reveals more insights into (a) the effects of purification protocols on the structure and function of isolated Stx2a (similar considerations may also be applicable to other toxins and proteins in general) and (b) the structure-function mechanisms underlying key points of eHUS pathogenesis such as Stx2a-binding to neutrophils, leukocyte-platelet aggregate formation, and FH binding abilities.

Different Stx purifications utilizing different bacterial strains and different purification protocols lead to different findings using purified Stx2a. On the other hand, based on the structure/function relationship described here, an important goal would be the characterisation of the structure of Stx2a circulating in EHEC-infected patients and its role in the pathogenetic process, that is, development of eHUS or recovery after the intestinal prodromal phase. However, a formal evidence of the presence of the cleaved toxin in patients’ circulation during EHEC infections is lacking. Work in progress in our laboratories is aimed at the specific detection of cleaved and uncleaved toxin forms in pathophysiological models of HUS.

Understanding the real structure and conformation of eHUS-inducing Stx2a would allow development of appropriate animal models aimed at studying the impact of the different toxin form (cleaved or uncleaved) on eHUS pathogenesis. This may improve strategies for the prevention of the life-threatening transition from hemorrhagic colitis to eHUS in EHEC-infected children.

4 | EXPERIMENTAL PROCEDURES

4.1 | Proteins and antibodies

LPS from E. coli serotype O111B4 was obtained from Alexis Biochemicals (Roma, Italy); FH from Calbiochem (Darmstadt, Germany); bovine serum albumin (BSA), thrombin, trypsin type XI (from bovine pancreas (T1005)), furin (F2677, 2000 U/ml), the furin inhibitor hexaarginine (SCP0148), 4-nitrophenyl phosphate disodium salt hexahydrate, and alkaline phosphatase (AP)-conjugated anti-sheep antibody (Ab) from Sigma Aldrich (Taufkirchen, Germany, or Milan, Italy); sheep anti-human FH (Ab) from Binding Site (Birmingham, UK); mouse anti-Stx2a Ab (Stx2-BB12) from Toxin Technology (Sarasota, FL); allophycocyanin (APC)-conjugated CD42b Ab from BD Biosciences (San Jose, CA); phycoerythrin (PE)-conjugated CD62P Ab from BioLegend (San Diego, CA); FITC-labelled anti-CD16, anti-CD65, anti-CD14, goat anti-mouse IgG, PE-labelled anti-CD41, PCS-labelled anti-CD16 from Beckman Coulter (Miami, FL); Blood (3.2% citrate) for the preparation of platelet-rich plasma (PRP) or serum were collected from three healthy volunteers after informed consent and with an existing ethical vote.

4.2 | Toxin purification

E. coli strain R82 (pJES) 120DH5a and E. coli C600 (933 W) were supplied by Mohamed Karmali (Ottawa, Canada) and by Alison O’Brien (Bethesda, MD), respectively. In the former case, Stx2a (named Stx2a(cl) throughout the paper) was purified in Innsbruck (Austria) by the following steps: culture, sonication, concentration on Amicon filters, dialysis, hydroxyapatite chromatography, dialysis, concentration on Amicon filters, chromatofocusing using DEAE Sepharose fast flow ion exchanger (Logan, Lagerlund, & Chamow, 1999), ammonium sulfate precipitation, and dialysis. In the second case, Stx2a (called Stx2a(uncI) throughout the paper) was isolated in Bologna (Italy) as described in a previous study (Matussek et al., 2003): culture, ammonium sulfate precipitation, dialysis, receptor analog affinity chromatography on Gala1-4Galβ-O-spacer-BSA-Sepharose 4B (Sterogene Bioseparations, Carlsbad, CA) to remove trace endotoxin contaminants. The amount of contaminating LPS was determined by using the Endosafe®Endochrome-K™ assay (Charles River, Charleston, SC). The test method is based on the Chromogenic technique as described elsewhere (United States Pharmacopeial Convention, Bacterial Endotoxins Test). Toxins were analysed by SDS-PAGE in reducing and non-reducing conditions using a 16% (wt/vol) gel followed by staining with Coomassie blue.

4.3 | Cleavage of Stxs with trypsin or furin

Stx2a(uncI) (4 μg) was incubated in 10 μl PBS pH 7.5 for 1 hr at 37°C with 50 ng of trypsin (1 mg/ml in 0.1 mM HCl, diluted to 0.05 ng/ml with PBS). Then, 0.7 ng of the trypsin inhibitor PMSF (1 mg/ml in absolute ethanol diluted to 0.7 μg/ml with water) was added for 10 min at 37°C. Alternatively, Stx2a(uncI; 4 μg) was incubated with 2 U of furin in 10 μl PBS pH 7.5 containing 1 mM CaCl₂ for 1 hr at 37°C. Then, 4 μl of the furin inhibitor hexa-D-arginine 1.05 mM (Cameron, Appel, Houghten, & Lindberg, 2000) was added for 10 min at 37°C. The test method is based on the Chromogenic technique as described elsewhere (United States Pharmacopeial Convention, Bacterial Endotoxins Test). Toxins were analysed by SDS-PAGE in reducing and non-reducing conditions using a 16% (wt/vol) gel followed by staining with Coomassie blue.

4.4 | Sequencing of stx2a gene

The stx2a subunit A and B genes were amplified using primers and PCR conditions described elsewhere (Orth et al., 2007). Purified PCR products were analysed on the Applied Biosystems 3500 Genetic Analyser according to standard Sanger sequencing procedures and according to the manufacturer’s protocol. Sequencing data were processed and analysed with Sequencing 5.2 Analysis Software (Applied Biosystems). Sequences were aligned using Serial Cloner Software Version 2.6.

4.5 | Isolation of human neutrophils and binding assays with toxins

Neutrophils (99.7% purity) were isolated under endotoxin low conditions from buffy coats of several healthy donors by centrifugation over Ficoll-Paque, dextran sedimentation, erythrocytic lysis, and negative selection of contaminant cells (Stemcell Technologies, Vancouver, BC, Canada) as in the previous study (Brigotti et al., 2013; Cassatella et al., 1988). For binding experiments with Stx2a, Eppendorf tubes
were precoated with PBS containing 1% endotoxin low (≤1 endotoxin unit/mg) BSA to avoid nonspecific loss of the toxins (van Setten, Monners, Verstraten, van den Heuvel, & van Hinsbergh, 1996). Neutrophils (5 × 10⁵) were incubated 90 min at 37°C with different concentrations of Stx2a (2–60 nM) in 250 μl of the same buffer (PBS-BSA). After incubation, the cells were collected by centrifugation at 200 × g for 5 min and washed three times. Stx2a bound to neutrophils was detected by flow cytometry using a mouse monoclonal anti-Stx2a antibody as previously described. Binding of Stx2a to cells was quantified as mean channel value of fluorescence (Tazzari et al., 2004). The presence of the complexes trypsin/PMSF or furin/hexa-D-arginine in neutrophil-binding assays did not affect the binding activity of Stx2a(uncl). Cleaved or uncleaved Stx2a were similarly recognised (Raji assay, see below) by the B chain specific monoclonal antibody (Figure S3).

4.6 | Neutrophil-binding assay with Stx2a in human blood

Blood (250 μl) from healthy donors was challenged with Stx2a (60 nM) for 90 min at 37°C. Cells were collected by centrifugation at 2700 × g at 15 min at room temperature. After erythrocytic lysis, washed leukocytes were obtained, and the binding of Stx2a to neutrophils was assessed by indirect flow cytometric analysis as described above.

4.7 | Detection of leukocyte/platelet aggregates by direct flow cytometry

Whole-blood samples (1 ml) from various donors were incubated for 4 hr at 37°C with Stx2a (1 nM). After erythrocytic lysis, samples were incubated with PE-labelled anti-CD41 to reveal platelets and FITC-labelled anti-CD14 for monocyte detection or PC5-labelled anti-CD16 to identify neutrophils on gated granular cells. Controls with appropriate isotype Abs were performed. CD14/CD41 double-positive cell populations or CD16/CD41 granular double-positive cell populations were identified as monocyte/platelet or neutrophil/platelet aggregates, respectively.

4.8 | Analysis of Stx2a binding to FH

Binding of Stx2a to FH was detected by ELISA as previously described (Orth et al., 2009). The different Stx2a preparations were immobilised at 2 μg/well each in a microtiter plate followed by addition of FH at 2 μg/well for 6 hr at 37°C.

4.9 | Platelet activation assay

To evaluate whether Stx2a enhanced platelet activity, PRP was incubated for 90 min at 37°C with various concentrations of purified Stx2a (1–100 ng/ml). For positive control, PRP was treated with thrombin (0.1 U/ml), for negative control with medium only. Platelet activation and subsequent secretion of α-granules were evaluated via CD62P (P-selectin) quantification using FACScan apparatus (Becton Dickinson, NJ; Lu & Malinauskas, 2011). The antigen CD42b, present on both resting and activated platelets, was used for platelet identification (H. Suzuki, Yamamoto, Tanoue, & Yamazaki, 1987). Results were given as the mean fluorescence intensity.

4.10 | Radioactive translation inhibition assay with Raji cells

Inhibition of protein synthesis by Stx2a was assayed as previously described (Arfilli et al., 2015). Briefly, after 3-hr incubation of Raji cells (0.3 × 10⁶, supplied by Andrea Bolognesi, Bologna, Italy) with toxins (16.7 M–3.125 nM) in the presence or absence of 10% human sera, [3H]leucine (2 μCi) was added. Radioactive proteins synthesised during 1-hr incubation were collected on glass filters after treatment of cell pellets with KOH 0.1 N followed by 10% trichloroacetic acid (TCA) precipitation. After extensive washing with 5% TCA, the radioactivity was measured by a liquid-scintillation β counter.

4.11 | Vero cells cytotoxicity assay

The cytotoxicity assay using Vero cells was performed as previously described (Gentry & Dalrymple, 1980; Russo, Melton-Celsa, Smith, & O’Brien, 2014). Briefly, cells were incubated with toxins (150 aM–150 nM) for 48 hr. Detection was done via 1% crystal violet staining.

4.12 | Photophysical measurements

Absorption spectra were recorded on a Perkin Elmer l650 spectrophotometer. CD spectra were obtained with a Jasco J-715 spectropolarimeter (2 nm resolution) and fluorescence spectra on an Edinburgh spectrofluorimeter with right angle detection geometry at 22°C. A time-correlated single photon counting system (IBH Consultants Ltd.) was used to study the time-resolved fluorescence of air-equilibrated solutions, excited at 278 nm with a Nano-LED source. Decay profiles were collected at 330 nm and analysed using a multiexponential function as described (Brigotti et al., 2011; Lakowicz, 2006).

4.13 | Statistical analyses

Statistical analysis was performed with GraphPad Prism 5. Differences in continuous variables were tested with a t test after controlling the normality of their distribution. A P value <0.05 was considered statistically significant. Correlation between variables was assessed using a Pearson correlation coefficient.

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REFERENCES

Arfilli, V., Carnicelli, D., Ardissino, G., Torresani, E., Scavia, G., & Brigotti, M. (2015). A rapid and sensitive method to measure the functional activity of Shiga toxins in human serum. Toxins (Basel), 7, 4564–4576. https://doi.org/10.3390/toxins7114564

Armstrong, G. D., Mulvey, G. L., Marcato, P., Griener, T. P., Kahan, M. C., Tennent, G. A., ... Pepys, M. B. (2006). Human serum amyloid P component protects against Escherichia coli O157:H7 Shiga toxin 2 in vivo: Therapeutic implications for hemolytic-uremic syndrome. The Journal of Infectious Diseases, 193, 1120–1124. https://doi.org/10.1086/501472

Bauwens, A., Betz, J., Meisen, I., Kemper, B., Karch, H., & Muthing, J. (2013). Facing glycosphingolipid-Shiga toxin interaction: Dire straits for endothelial cells of the human vasculature. Cellular and Molecular Life Sciences, 70, 425–457. https://doi.org/10.1007/s00018-012-1060-z

Bauwens, A., Bielaszewska, M., Kemper, B., Langehanenberg, P., von Bally, G., Reichelt, R., ... Muthing, J. (2011). Differential cytotoxic actions of Shiga toxin 1 and Shiga toxin 2 on microvascular and macrovascular endothelial cells. Thrombosis and Haemostasis, 105, 515–528. https://doi.org/10.1160/TH10-02-0140

Bergan, J., Dyve Lingelem, A. B., Skotland, T., & Sandvig, K. (2012). Shiga toxins. Toxicon, 60, 1085–1107. https://doi.org/10.1016/j.toxicon.2012.07.016

Brahms, S., & BrahmS, J. (1980). Determination of protein secondary structure by solution of vacum ultraviolet circular dichroism. Journal of Molecular Biology, 138(2), 149–178. https://doi.org/10.1016/0022-2836(80)90282-X

Brigotti, M., Alfieri, R., Sestili, P., Bonelli, M., Petronini, P. G., Guidarelli, A., ... Serti, P. (2002). Damage to nuclear DNA induced by Shiga toxin 1 and ricin in human endothelial cells. The FASEB Journal, 16(3), 365–372. https://doi.org/10.1096/fj.01-0521com

Brigotti, M., Carnicelli, D., Arfilli, V., Rocchi, L., Ricci, F., Pagliaro, P., ... Monti, S. (2011). Change in conformation with reduction of alpha-helix content causes loss of neutrophil binding activity in fully cytotoxic Shiga toxin 1. The Journal of Biological Chemistry, 286(40), 34514–34521. https://doi.org/10.1074/jbc.M111.255414

Brigotti, M., Carnicelli, D., Arfilli, V., Tamassia, N., Borsetti, F., Fabbrì, E., ... Cassatella, M. A. (2013). Identification of TLR4 as the receptor that recognizes Shiga toxins in human neutrophils. The Journal of Immunology, 191, 4748–4758. https://doi.org/10.4049/jimmunol.1300122

Brigotti, M., Carnicelli, D., Ravaneli, E., Barbieri, S., Ricci, F., Bontadini, A., ... Tazzari, P. L. (2008). Interactions between Shiga toxins and human polymorphonuclear leukocytes. Journal of Leukocyte Biology, 84, 1019–1027. https://doi.org/10.1189/jlb.0308157

Brigotti, M., Carnicelli, D., Ravaneli, E., Varà, A. G., Martinelli, C., Alfieri, R. R., ... Sestili, P. (2007). Molecular damage and induction of proinflammatory cytokines in human endothelial cells exposed to Shiga toxin 1. Shiga toxin 2, and alpha-sarcin. Infection and Immunity, 75, 2201–2207. https://doi.org/10.1128/IAI.01707-06

Cameron, A., Appel, J., Houghten, R. A., & Lindberg, I. (2000). Polyarginines are potent furin inhibitors. The Journal of Biological Chemistry, 275, 36741–36749. https://doi.org/10.1074/jbc.M003848200

Cassatella, M. A., Cappelli, R., Della Bianca, V., Grzeskowiak, M., Dusi, S., & Berton, G. (1988). Interferon-gamma activates human neutrophil oxygen metabolism and exocytosis. Immunology, 63, 499–506.
Matussek, A., Lauber, J., Bergau, A., Hansen, W., Rohde, M., Dittmar, K. E., Gunzer, F. (2003). Molecular and functional analysis of Shiga toxin-induced response patterns in human vascular endothelial cells. Blood, 102, 1323–1332. https://doi.org/10.1182/blood-2002-10-3301

Melton-Celsa, A. R., Darnell, S. C., & O’Brien, A. D. (1996). Activation of Shiga-like toxins by mouse and human intestinal mucus correlates with virulence of enterohemorrhagic Escherichia coli O91:H21 isolates in orally infected, streptomycin-treated mice. Infection and Immunity, 64, 1569–1576.

Obrig, T. G. (2010). Escherichia coli Shiga toxin mechanisms of action in renal disease. Toxins (Basel), 2, 2769–2794. https://doi.org/10.3390/toxins2122769

Orth, D., Grif, K., Khan, A. B., Naim, A., Dierich, M. P., & Wurzner, R. (2007). The Shiga toxin genotype rather than the amount of Shiga toxin or the cytotoxicity of Shiga toxin in vitro correlates with the appearance of the hemolytic uremic syndrome. Diagnostic Microbiology and Infectious Disease, 59, 235–242. https://doi.org/10.1016/j.diagmicrobio.2007.04.013

Orth, D., Khan, A. B., Naim, A., Grif, K., Brockmeyer, J., Karch, H., ... Wurzner, R. (2009). Shiga toxin activates complement and binds factor H: Evidence for an active role of complement in hemolytic uremic syndrome. The Journal of Immunology, 182, 6394–6400. https://doi.org/10.4049/jimmunol.0900151

Poolpol, K. (2012). PhD Dissertation. (PhD). Austria: Medical University of Innsbruck.

Poolpol, K., Orth-Holler, D., Speth, C., Zipfel, P. F., Skerka, C., de Cordoba, S. R., ... Wurzner, R. (2014). Interaction of Shiga toxin 2 with complement regulators of the factor H protein family. Molecular Immunology, 58, 77–84. https://doi.org/10.1016/j.molimm.2013.11.009

Rosales, A., Hofer, J., Zimmerhackl, L. B., Jungrathmayr, T. C., Riedl, M., Giner, T., ... for the German-Austrian HUS Study Group (2012). Need for long-term follow-up in enterohemorrhagic Escherichia coli-associated hemolytic uremic syndrome due to late-emerging sequelae. Clinical Infectious Diseases, 54, 1413–1421. https://doi.org/10.1093/cid/cis196

Russo, L. M., Melton-Celsa, A. R., Smith, M. J., & O’Brien, A. D. (2014). Comparisons of native Shiga toxins (Sxs) type 1 and 2 with chimeric toxins indicate that the source of the binding subunit dictates degree of toxicity. PLoS One, 9, e93463. https://doi.org/10.1371/journal.pone.0093463

Sandvig, K., Garred, O., Prydz, K., Kozlov, J. V., Hansen, S. H., & van Deurs, B. (1992). Retrograde transport of endocytosed Shiga toxin to the endoplasmic reticulum. Nature, 358, 510–512. https://doi.org/10.1038/358510a0

Sandvig, K., & van Deurs, B. (2005). Delivery into cells: lessons learned from plant and bacterial toxins. Gene Therapy, 12, 865–872. https://doi.org/10.1038/sj.gt.3302525

Stahl, A. L., Sartz, L., & Karpman, D. (2011). Complement activation on platelet-leukocyte complexes and microparticles in enterohemorrhagic Escherichia coli-induced hemolytic uremic syndrome. Blood, 117, 5503–5513. https://doi.org/10.1182/blood-2010-09-309161

Stahl, A. L., Sartz, L., Nilsson, A., Bekassy, Z. D., & Karpman, D. (2009). Shiga toxin and lipopolysaccharide induce platelet-leukocyte aggregates and tissue factor release, a thrombotic mechanism in hemolytic uremic syndrome. PLoS One, 4, e6990. https://doi.org/10.1371/journal.pone.0006990

Suzuki, A., Doi, H., Matsuzawa, F., Akawa, S., Takiguchi, K., Kawano, H., ... Ohno, S. (2000). Bcl-2 antiapoptotic protein mediates verotoxin II-induced cell death: possible association between bcl-2 and tissue failure by E. coli O157:H7. Genes & Development, 14, 1734–1740.

Suzuki, H., Yamamoto, N., Tanoue, K., & Yamazaki, H. (1987). Glycoprotein Ib distribution on the surface of platelets in resting and activation states: An electron microscope study. Teihe Histochemical Journal, 19, 125–136. https://doi.org/10.1007/BF01695137

Tarr, P. I., Gordon, C. A., & Chandler, W. L. (2005). Shiga-toxin-producing Escherichia coli and haemolytic uraemic syndrome. Lancet, 365, 1073–1086. https://doi.org/10.1016/S0140-6736(05)11444-2

Tazzari, P. L., Ricci, F., Carnicelli, D., Caprioli, A., Tozzi, A. E., Rizzoni, G., ... Brigotti, M. (2004). Flow cytometry detection of Shiga toxins in the blood from children with hemolytic uremic syndrome. Cytometry Part B Clinical Cytometry, 61, 40–44.

van Setten, P. A., Monnens, L. A., Verstraten, R. G., van den Heuvel, L. P., & van Hinsbergh, V. W. (1996). Effects of verocytotoxin-1 on nonadherent human monocytes: Binding characteristics, protein synthesis, and induction of cytokine release. Blood, 88, 174–183.

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