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RESEARCH ARTICLE

Novel IgG-Degrading Enzymes of the IgdE Protease Family Link Substrate Specificity to Host Tropism of Streptococcus Species

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Abstract

Recently we have discovered an IgG degrading enzyme of the endemic pig pathogen S. suis designated IgdE that is highly specific for porcine IgG. This protease is the founding member of a novel cysteine protease family assigned C113 in the MEROPS peptidase database. Bioinformatical analyses revealed putative members of the IgdE protease family in eight other Streptococcus species. The genes of the putative IgdE family proteases of S. agalactiae, S. porcinus, S. pseudoporcinus and S. equi subsp. zooepidemicus were cloned for production of recombinant protein into expression vectors. Recombinant proteins of all four IgdE family proteases were proteolytically active against IgG of the respective Streptococcus species hosts, but not against IgG from other tested species or other classes of immunoglobulins, thereby linking the substrate specificity to the known host tropism. The novel IgdE family proteases of S. agalactiae, S. pseudoporcinus and S. equi showed IgG subtype specificity, i.e. IgdE from S. agalactiae and S. pseudoporcinus cleaved human IgG1, while IgdE from S. equi was subtype specific for equine IgG7. Porcine IgG subtype specificities of the IgdE family proteases of S. porcinus and S. pseudoporcinus remain to be determined. Cleavage of porcine IgG by IgdE of S. pseudoporcinus is suggested to be an evolutionary remaining activity reflecting ancestry of the human pathogen to the porcine pathogen S. porcinus. The IgG subtype specificity of bacterial proteases indicates the special importance of these IgG subtypes in counteracting infection or colonization and opportunistic streptococci neutralize such antibodies through expression of IgdE family proteases as putative immune evasion factors. We suggest that IgdE family proteases might be valid vaccine targets against streptococci of both human and veterinary medical concerns and could also be of therapeutic as well as biotechnological use.
Competing Interests: We have read the journal’s policy and the authors of this manuscript have the following competing interests: A patent application [New streptococcal proteases, 1630021-2] for IgdE proteases has been filed. CS and UPR are listed as inventors in this application. This does not alter our adherence to PLOS ONE policies on sharing data and materials.

Introduction

Gram-positive bacteria of the genus *Streptococcus* are highly intertwined with humans and animals as commensal, opportunistic and pathogenic bacteria. Often streptococci show pronounced host tropism, but these bacteria can also cause zoonotic or anthroponotic infections in more uncommon hosts [1].

Starting with the observation of a Immunoglobulin (Ig) G degrading activity in culture supernatants of the important endemic pig pathogen *Streptococcus (S.) suis*, we identified recently a novel IgG degrading enzyme designated IgdE [2]. Through inhibitor screening, *in silico* modeling and mutational studies of the potential catalytic triad residues, IgdE was assigned to be a cysteine protease. This protease does not have homology to any protease previously described and is thereby the founding member of a novel cysteine protease family designated C113 within the CA clan in the MEROPS peptidase database (https://merops.sanger.ac.uk/) [3]. IgdE of *S. suis* is highly specific for porcine IgG and no other substrate has been identified. Immune evasion from Ig mediated immune defense seems to be of special importance for bacteria of the *Streptococcus* genus. Several Ig degrading enzymes of streptococci have been identified, such as the IgG specific proteases, IdeS of *S. pyogenes* [4], IdeZ of *S. equi* subsp. *zooepidemicus* and IdeE of *S. equi* subsp. *equi* [5]. In addition IgA-specific proteases of *S. pneumoniae*, *S. oralis*, *S. sanguis* and *S. mitis* have been described [6, 7] and recently, we described an IgM specific protease in *S. suis*, designated IdeSsuis [8]. Most of these proteases cleave the heavy chain of Ig molecules in the hinge region, thereby impairing all effector functions of Ig except neutralization.

IgG is the major antibody in serum of most higher organisms [9, 10]. Human IgG is categorized into four different subclasses; IgG1, IgG2, IgG3 and IgG4, with abundance in the same order [11]. These subclasses have subtle variations in structure resulting in different effector mechanisms. IgG1 and IgG3 are the prevalent Ig classes in humans able to cross the placental barrier to protect the fetus and newborn by passive immunization [12]. Every IgG subclass has an individual FcγR-binding profile with IgG1 and IgG3 binding efficiently to most FcγR, while IgG2 and IgG4 have a reduced affinity for some FcγR [13]. IgG1 and IgG3 activate the classical complement pathway efficiently compared to IgG2 and IgG4 [14–16]. Consequently IgG1 and IgG3 are more involved in complement activation, phagocyte binding, sensitization of mast cells and sensitization of natural killer cells for killing, while all four subclasses are equally involved in neutralization, opsonization and extravascular diffusion [17].

Porcine IgG is divided into six subclasses [18]. These subclasses are considered to be biochemically inseparable. Thus the different IgG subtypes and their proposed properties are only predicted by sequence analyses [19]. Equine IgG is divided into seven subclasses [20]. Experimental studies with recombinant equine IgG subclasses revealed that IgG1, IgG3, IgG4 and IgG7 are the most potent activators of the classical complement pathway via C1q binding and elicit also a strong respiratory burst from equine peripheral blood leukocytes [21].

In this study, we employed rigorous homology searches to identify several homologues of IgdE of *S. suis* in other *Streptococcus* species as putative IgdE family proteases and compared them by phylogenetic analysis. The *igdE* genes of *S. agalactiae*, *S. porcinus*, *S. pseudoporicinus* and *S. equi* subsp. *zooepidemicus* were cloned for expression and purification of recombinant protein followed by screening for potential substrates of these putative proteases.

*S. agalactiae*, also known as Group B *Streptococcus*, is commonly found as a commensal in cattle and humans [22, 23], but is also able to cause mastitis in cows [24] and genitourinary infections, neonatal sepsis, CNS infections and endocarditis in humans [25].

*S. porcinus* is a bacterium most commonly found in the respiratory tract of pigs [26] and has been associated with lymphadenitis [27] and still birth [28]. *S. pseudoporicinus* has recently
been distinguished from *S. porcinus* as a separate species [29] and has been shown to be an emerging and common organism colonizing the genitourinary tract of women [30].

*S. equi* subsp. *zooepidemicus*, a commensal and opportunistic pathogen of horses as well as other mammals, can cause severe zoonotic infections in humans, such as sepsis, meningitis and endocarditis [31]. In horses *S. equi* subsp. *zooepidemicus* can cause several different pathologies, including respiratory tract infections, uteritis and wound infections. The cause of the highly contagious upper respiratory tract disease strangles, *S. equi* subsp. *equi*, is believed to be a clonal descendant of an ancestral strain of *S. equi* subsp. *zooepidemicus* [32, 33].

All putative IgdE family proteases tested in this study showed enzymatic activity and substrate specificity for IgG of specific hosts. IgdE of *S. agalactiae* was specific for human IgG1. IgdE of *S. porcinus* was specific for porcine IgG, while IgdE of *S. equi* subsp. *zooepidemicus* was specific for equine IgG7. IgdE of *S. pseudoporcinus* degraded both human IgG1 as well as porcine IgG thereby being the only identified IgdE family protease with multiple substrates. The substrate specificities, in regard of host species IgG, of these novel members of the IgdE protease family correlate well with the known host tropism of the respective *Streptococcus* species. The IgG subclass specificities of these proteases might implicate special importance of these specific IgG subtypes in immune defense against these *Streptococcus* species during certain stages of infection which the bacteria might counteract through expression of IgG subtype specific proteases.

The IgdE proteases of *S. agalactiae*, *S. porcinus*, *S. pseudoporcinus* and *S. equi* subsp. *zooepidemicus* are designated IgdE$_{agalactiae}$, IgdE$_{porcinus}$, IgdE$_{pseudoporcinus}$ and IgdE$_{equi}$ in this study.

**Materials and Methods**

**Computational identification of novel IgdE protease family members within Streptococci**

Coding sequences of all available *Streptococcus* genomes were downloaded from NCBI ([ftp://ftp.ncbi.nlm.nih.gov/genomes/Bacteria/](ftp://ftp.ncbi.nlm.nih.gov/genomes/Bacteria/) on Aug-21-2015) and from PATRIC ([ftp://ftp.patricbrc.org/patric2/](ftp://ftp.patricbrc.org/patric2/) on Aug-25-2015). As a reference sequence for an IgdE protease the RefSeq sequence WP_014636499.1 of *S. suis* was used. The N-terminal signal peptide and the C-terminal region only present in sequences from *S. suis* were removed, leaving amino acids 38–520, hereafter called IgdE$_{domain}$.

The IgdE$_{domain}$ was used as query sequence in blastp searches (E-value cutoff 1 to keep all possible proteases) against the NCBI sequences as well as the PATRIC sequences. Sequences not containing the catalytic cysteine were removed from further consideration. Many of the sequences found are annotated as S-layer proteins or as containing an S-layer homology domain W. These are often present in two or more copies in the same genome, and have an SxC or GxC motif in the catalytic site instead of the AxC motif found in aa 300–302 of the original IgdE sequence of *S. suis*. In order to distinguish these sequences, which are not members of the IgdE protease family, all sequences lacking the AxC motif were also removed.

The obtained hits were in turn used as query sequences against the same databases, using the same parameters. From the list of matched sequences those that in the second round had a match overlapping with the region matched in the first round, when the IgdE$_{domain}$ was used as query, were chosen. Sequences matching to the Transglutcore model with an E-value of at most 1e-6 or sequences not containing the catalytic AxC motif were excluded. The remaining sequences were trimmed at both ends to contain only the parts matching the IgdE$_{domain}$ sequence. In cases where this resulted in identical sequences only one copy was kept.
Phylogenetic analysis

Clustal Omega version 1.2.1 (http://www.ebi.ac.uk/Tools/msa/clustalo/) [34] was used to generate a multiple sequence alignment of the sequences obtained above using default settings. To determine the best fitting amino acid substitution model we used ProtTest version 3.4 [35]. A Jones-Taylor-Thornton (JTT) model with a gamma distribution, a proportion of invariable sites, and observed amino acid frequencies was the best model, and therefore used to construct a maximum likelihood (ML) tree with PhyML version 20131022 [36]. To assess the significance of phylogenetic grouping a bootstrap analysis with 100 repetitions was performed. The tree was rooted using an out-group consisting of two non-streptococcal protein sequences homologous to the IgdE_domain (WP_029500965.1; WP_016310821.1). These sequences were trimmed at both ends to contain only the parts matching the IgdE_domain sequence. Encoded proteins lack described functions. The phylogenetic tree was visualized with iTOL (http://itol.embl.de/) [37].

Bacterial strains and growth conditions

*Escherichia coli* strains were cultured in Lysogeny Broth (LB) or Lysogeny Agar (LA) under aerobic conditions at 30°C or 37°C. When appropriate, 50 μg/ml kanamycin or 25 μg/ml chloramphenicol was added.

Cloning of IgdE homologues for recombinant expression

Genes of the IgdE homologues lacking the signal peptide were amplified from chromosomal DNA of *S. porcinus* strain DSM20725 (kindly received from Christoph G. Bauns, College of Veterinary Medicine, University Leipzig, Leipzig, Germany), *S. pseudoporcinus* strain LQ940-04T (ATCC), *S. agalactiae* strain CCUG420 (kindly received from Åsa Gylfe, Department of Clinical Microbiology, Umeå University, Umeå, Sweden) and *S. equi* subsp. *zooepidemicus* strain 203 (kindly received from National Veterinary Institute, Uppsala, Sweden) as templates using primer pairs designated in Table 1. PCR fragments were cloned into pET_ZZ_1a after digestion with restriction enzymes (all Thermo Scientific) denoted in primer names. The cloned plasmids were verified by sequencing and transformed into *E. coli* BL21 (DE3) pLysS for recombinant expression of the proteins.

Expression and purification of recombinant IgdE homologues

*E. coli* BL21(DE3) pLysS isolates carrying pET_ZZ_1a igdE*porcinus*, igdE*pseudoporcinus*, igdE*agalactiae* or igdE*equi* were grown to OD600 0.5 at 30°C. Protein expression was induced with 0.5 mM IPTG for 5h at 30°C. Cells were lysed for crude soluble extracts by BugBuster HT Protein Extraction Reagent (Novagen) according to manufacture protocol or lysed by sonication in 20 mM sodium phosphate, 0.5M NaCl, 40 mM imidazole, pH 7.4 prior to further purification. The His-ZZ-tagged proteins were purified on HisTrap FF (GE Healthcare) using standard protocols. The tag was removed by enzymatic cleavage by Tev-protease for 20h at 4°C followed by a second round of purification on HisTrap FF (GE Healthcare). The flow through, containing untagged recombinant protein, was collected and buffer exchanged against PBS. Protein concentrations were determined by Nanodrop A280 measurements at appropriate dilutions. In case of IgdE*equi* no great overexpression was achieved and no purification attempt was conducted.

Screening for Ig-degrading activities of recombinant IgdE homologues

If not stated otherwise, all reactions were carried out at 37°C for 16h in PBS. 20 μg/ml purified recombinant proteins or 5% crude soluble extracts of *E. coli* expressing the igdE constructs...
were incubated with 0.5 mg/ml porcine, human, bovine, horse, goat, and mouse IgG (all Sigma), 0.25 mg/ml human IgG1 kappa, IgG2 kappa, IgG3 kappa, IgG4 kappa, IgA and IgM (all Sigma), 0.09 mg/ml purified recombinant horse IgG1, IgG2, IgG3, IgG4, IgG5, IgG6 and IgG7 expressed in Chinese hamster ovary cells or FreeStyle 293-F cells (according to [21]) and 1% human plasma, porcine plasma (kindly received from Christoph G. Baums, College of Veterinary Medicine, University Leipzig, Leipzig, Germany) and equine serum (Sigma). Reaction samples were analyzed using SDS-PAGE or Western Blot analyses. Experiments were repeated at least two times and representative analyses are shown.

SDS-PAGE and Western Blot analysis

Samples for SDS-PAGE were prepared with reducing sample buffer and heated to 95°C for 5 min. 12% SDS-PAGE was either stained with Coomassie blue (Sigma), Coomassie Fluor™ Orange Protein Gel Stain (Invitrogen) or blotted to Hybond-P PVDF membrane (GE Healthcare) for Western Blot analyses. Membranes were blocked with 5% dry milk powder in 0.1% PBS-Tween, followed by incubation with horse-radish peroxidase conjugated primary antibodies or unconjugated primary antibody (according to Table 2). Membranes were thoroughly washed with 0.1% PBS-Tween and development with Amersham ECL Select Western blotting detection reagent (GE Healthcare) according to manufacturer’s instruction and chemiluminescent signal was captured by LAS4000 imaging system (Fujifilm). Prestained protein ladders were pictured with the same system by standard epi-white illumination.

N-terminal Edman sequencing

IgG degradation reactions were separated by SDS-PAGE as previously explained and transferred to PVDF blotting membrane (GE Health Care) by semi-dry blotting with blotting buffer consisting of 50 mM Sodium borate and 20% Methanol. The membrane was stained with

Table 1. Used primers for cloning of igdE genes.

| Locus tag       | aa   | primers                                      |
|-----------------|------|---------------------------------------------|
| igdE<sub>porcinus</sub> | 34–527 | igdE<sub>porcinus</sub>-frw_NcoI, igdE<sub>porcinus</sub>-rev_Acc65l |
| STRPO_RS07810  |      | GTACCCATGGCTGTCTCTGGAGACAGAAATAG             |
|                 |      | GTACGGATCTTTAGTTATCCACATTCTTTTGTTC          |
| igdE<sub>pseudoporcinus</sub> | 38–535 | igdE<sub>pseudoporcinus</sub>-frw_Eco311, igdE<sub>pseudoporcinus</sub>-rev_Acc65l |
| STRPS_RS03610  |      | GTACGTTCTCCATGAGAAATAATATTTAAGAAAAACGTAAGCAATTAC |
|                 |      | GTACGGTACCTTACTGTCAGCTTTGTTTGAG             |
| igdE<sub>agalactiae</sub> | 37–623 | igdE<sub>agalactiae</sub>-frw_BspHI, igdE<sub>agalactiae</sub>-rev_Acc65l |
| MSA_19930       |      | GTACTCATGAACTAAATATTTTAAAGAAAAACGTAAGCAATTAC |
|                 |      | GTACGGTACCTTATCGTTATTTGCTTC                |
| igdE<sub>equi</sub> | 1-517(no signal peptide) | igdE<sub>equi</sub>-frw_NcoI, igdE<sub>equi</sub>-rev_Acc65l |
| M837_01916      |      | GTACCCATGGAGACATGAAAGAGTAG                |
|                 |      | GTACGGTACCTTATCGATTGAGCGTCTTACATG         |

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Table 2. Used antibodies and dilutions for Western Blot analysis.

| Name                | Manufacturer             | Dilution |
|---------------------|--------------------------|----------|
| Goat anti-pig IgG-HRP | Thermo Scientific (PA1-28685) | 1:30'000 |
| Goat anti-pig IgM-HRP | Thermo Scientific (PA1-84622) | 1:30'000 |
| Goat anti-pig IgA-HRP | Thermo Scientific (PA1-84625) | 1:1'000  |
| Rabbit anti-horse IgG-HRP | Abcam (ab6921) | 1:30'000 |
| Goat anti-horse IgM-HRP | Abcam (ab112879) | 1:1'000  |
| Rabbit anti-horse IgA-HRP | Abcam (ab112871) | 1:1'000  |

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Ponceau S 0.5% (Sigma) 1% acetic acid in MQ water and destained with MQ water. The membrane was quickly dried, where after the degradation product was neatly cut out. N-terminal Edman sequencing of the degradation product was performed by Proteome Factory (Berlin, Germany). BLAST homology searches were used to identify the position of the obtained sequence in the IgG molecule.

Results
Putative IgdE family proteases are spread among several Streptococcus species

To identify novel IgdE family proteases, streptococcal genomes were searched for sequences encoding homologues of the IgdE_domain. The IgdE_domain was defined as amino acids 38–520 of the RefSeq sequence WP_014636499.1 of S. suis 05ZYH33, thereby excluding the N-terminal signal peptide and the C-terminal region only present in sequences from S. suis. Only sequences that, when used as query in a second round of homology searches, had an overlap with the region matched in the first round and carried the conserved catalytic cysteine residue were kept for further analysis (Table 3).

After homology searches and filtering steps, 55 unique sequences were identified representing putative IgdE family proteases, 23 from S. suis, 18 from S. agalactiae, five from S. dysgalactiae, three from S. equi subsp. zooepidemicus and one each from S. porcinus, S. pseudoporcinus, S. canis and S. castoreus respectively. Two sequences from the same genome of S. merionis were identified. All sequences from S. agalactiae are very similar in the defined IgdE_domain region, often only differing in one or a few amino acid positions.

To illustrate the relationship of these 55 sequences a phylogenetic tree was inferred by maximum likelihood based on the JTT model of sequence evolution (Fig 1). The IgdE_domain region sequences of S. canis and S. castoreus showed similarity to the IgdE_domain region sequences of S. dysgalactiae. The two IgdE_domain sequences of S. merionis grouped together, but were still distinct from each other. IgdE_domain region sequences of all S. agalactiae strains grouped close together, as did sequences of S. equi and S. dysgalactiae, while the sequences of S. suis were more diverse. Also the two sequences obtained from S. porcinus and S. pseudoporcinus grouped close together.

Locations of igdE genes within Streptococcus genomes

Each gene encoding a putative IgdE protease was localized in the respective genome as well as genes flanking the igdE gene (Table 4). Location of the igdE genes was generally conserved.
within species, although two out of 18 *S. agalactiae* strains and one out of 23 *S. suis* strains had deviant neighboring genes. Location and neighboring genes were conserved between the species *S. castoreus*, *S. canis* and *S. dysgalactiae*. For sequences retrieved from genome drafts the locations could not be determined or could only be approximated.

**IgdE family proteases are highly specific for IgG of different host species**

The *igdE* genes from one representative strain of *S. agalactiae*, *S. porcinus*, *S. pseudoporcinus* and *S. equi* subsp. *zooepidemicus* (corresponding to the sequences marked in **bold** in Fig 1) were cloned into expression vectors in *E. coli*. The encoded proteins were over-expressed and recombinant protein was used for substrate screening by overnight incubation with potential substrates prior to analyses by SDS-PAGE and western blots.

Recombinant IgdE<sub>agalactiae</sub> was able to degrade human IgG and a diagnostic cleavage product of 32 kDa appeared, when rIgdE<sub>agalactiae</sub> was incubated with human IgG, but not when incubated with porcine, bovine, equine, caprine or murine IgG (Fig 2A). Similar to that rIgdE<sub>porcinus</sub> was only able to degrade porcine IgG; the diagnostic cleavage product of 32 kDa appeared only when incubated with porcine IgG, but not when incubated with human, bovine, equine, caprine or murine IgG (Fig 2B). Recombinant IgdE<sub>pseudoporcinus</sub> had in contrast dual substrate specificity towards human and porcine IgG, again characterized through appearance of diagnostic cleavage products of 32 kDa, while no degradation of bovine, equine, caprine or murine IgG could be observed (Fig 2C). Recombinant IgdE<sub>equi</sub> possessed degrading activity against equine IgG, while no degradation of human, porcine, bovine, caprine or murine IgG could be observed (Fig 2D).

IgG specificity of these novel members of the IgdE protease family was investigated by incubation of recombinant protein with porcine and equine serum, respectively. These cleavage reactions were analyzed by anti-IgG, anti-IgM and anti-IgA western blots. Recombinant IgdE<sub>porcinus</sub> degraded porcine serum IgG, but not IgM or IgA (Fig 3A). A similar observation was made when serum was incubated with rIgdE<sub>pseudoporcinus</sub> (Fig 3B), which also was also specific for IgG. Recombinant IgdE<sub>equi</sub> cleaved equine serum IgG, but not IgM or IgA (Fig 3C).

Specificity of the IgdE proteases from *S. agalactiae* and *S. pseudoporcinus* for human IgG in comparison to human IgM and IgA was tested through incubation of recombinant protein with purified Ig (all Sigma) and analyzed by reducing SDS-PAGE. The findings showed that rIgdE<sub>agalactiae</sub> (Fig 4A) and rIgdE<sub>pseudoporcinus</sub> (Fig 4B) were specific for IgG and cleaved human IgG, but not human IgM or IgA.

**IgG subtype specificity of IgdE proteases of *S. agalactiae*, *S. pseudoporcinus* and *S. equi* subsp. *zooepidemicus***

Given the shown specificity of IgdE proteases for IgG of specific hosts, we were interested in IgG subtype specificity of these proteases. Recombinant IgdE<sub>agalactiae</sub> (Fig 5A) and rIgdE<sub>pseudoporcinus</sub> (Fig 5B) were therefore incubated with human IgG1, IgG2, IgG3 and IgG4 from myeloma source prior to reducing SDS-PAGE analysis. Both rIgdE<sub>agalactiae</sub> and rIgdE<sub>pseudoporcinus</sub> were strictly IgG1 specific, and no degradation of human IgG2, IgG3 and IgG4 was observed. The equine IgG subtype specificity of rIgdE<sub>equi</sub> was tested by incubation of protease preparations with purified recombinant equine IgG1, IgG2, IgG3, IgG4, IgG5, IgG6 and IgG7 (Fig 5C). Strikingly, pronounced subtype specificity was also observed for rIgdE<sub>equi</sub> and of all equine IgG subtypes only recombinant equine IgG7 was cleaved.

**Cleavage sites of IgdE family proteases**

To determine the exact cleavage site of the IgdE proteases within the respective substrate IgG molecules, the 32 kDa cleavage products generated by IgdE<sub>agalactiae</sub>, IgdE<sub>porcinus</sub>
Fig 1. Phylogenetic tree of identified putative IgdE proteases. Phylogenetic analysis of the IgdE_domain of 55 identified putative IgdE proteases. The maximum likelihood (ML) tree was constructed using a Jones-Taylor-Thornton (JTT) model with PhyML. Bootstrap values greater than or equal to 80% are shown. Putative IgdE protease sequences that corresponded to the genes cloned for expression of recombinant protein are marked in bold. Tree scale is given as average number of substitutions per site.

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IgdE<sub>pseudoporcinus</sub> and IgdE<sub>equi</sub> were subjected to N-terminal Edman sequencing. All obtained sequences corresponded to the hinge region of the respective IgG molecules (Fig 6A). IgdE<sub>agalactiae</sub> and IgdE<sub>pseudoporcinus</sub> cleave both human IgG1, but cleavage sites are not identical, being located two residues apart from each other. IgdE<sub>agalactiae</sub> cleaves the human IgG1 heavy chain two residues N-terminal of the putative homodimer disulfide bond cysteine residues; while all other cleavage sites of IgdE family proteases are located directly adjacent to the putative homodimer disulfide bond in IgG hinge regions. The cleavage sites identified in porcine IgG generated through cleavage with both IgdE<sub>porcinus</sub> and IgdE<sub>pseudoporcinus</sub> were identical and found in porcine IgG2, IgG4 and IgG6 (Fig 6B). Similar sequences can, however, also be found in IgG1 and IgG5. The sequence obtained by N-terminal Edman sequencing of the cleavage product of equine IgG generated by IgdE<sub>equi</sub> corresponded to the hinge region of equine IgG7. The amino acid sequences analogous to those adjacent to the cleavage site in equine IgG7 are different in all the other horse IgG subtypes, providing a rationale for the observed subtype specificity.

**Discussion**

Based on the identification of the founding member of a novel cysteine protease family, IgdE of S. <i>suis</i> [2], we identified several putative IgdE family proteases through homology searches within the genus <i>Streptococcus</i>. Locations of the genes of these putative proteases were conserved within, and to certain degree also between, different <i>Streptococcus</i> species suggesting that igdE genes are part of the core genome and not part of mobile elements (Table 4). One sequence of S. <i>suis</i> and two of S. <i>agalactiae</i> had, however, different neighboring genes than the other 22 sequences of S. <i>suis</i> and 16 of S. <i>agalactiae</i>, respectively. Putative IgdE family protease sequences were however only found in 55% of S. <i>agalactiae</i>, 46% of S. <i>equi</i> and 81% of S. <i>suis</i> genomes (Table 3). This could be due to the real absence of an igdE gene in some strains of these species or due to pore, incomplete or missannotated coding sequences of these genomes.
Fig 2. IgG host species specificity of IgdE family proteases. 0.5 mg/ml human, porcine, equine, bovine and murine IgG were incubated for 16h at 37°C with (A) 20 μg/ml purified rIgdE agalactiae, (B) 20 μg/ml purified rIgdE porcinus, (C) 5% soluble fraction of E. coli cells expressing rIgdE pseudoporcinus, (D) 5% soluble fraction of E. coli cells expressing rIgdE equi. PBS (A and B) or 5% soluble fraction of E. coli cells without recombinant construct (C and D) were used as negative controls (-). Reactions were analyzed by Coomassie blue SDS-PAGE under reducing conditions. Images of different SDS-PAGE run in parallel have been assembled into one figure. The diagnostic 32 kDa IgG cleavage product (*) appeared when rIgdE agalactiae was incubated with human IgG, rIgdE porcinus with porcine IgG, rIgdE pseudoporcinus with human IgG and porcine IgG, and rIgdE equi with equine IgG.

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Fig 3. IgdE family proteases are specific for IgG compared to IgM and IgA. 2% porcine plasma was incubated with (+) or without (-) 20 μg/ml purified rIgdEporcinus (A) or 5% soluble fraction of E. coli cells expressing rIgdEpseudoporcinus (B) respectively for 16h at 37˚C. 2% equine serum was incubated with 5% soluble fraction of E. coli cells expressing rIgdEequi (C) for 16h at 37˚C. The reactions were analyzed by anti-porcine or anti-equine IgG, IgM and IgA Western blots under reducing conditions. Only degradation products of IgG (*) could be observed.

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All investigated IgdE family proteases except IgdE<sub>pseudoporcinus</sub> showed only specificity towards IgG of one host species. The substrate specificity of these IgdE family proteases correlates well with the known host tropism of the respective Streptococcus species. However, IgdE<sub>agalactiae</sub> does not cleave bovine IgG, despite <i>S. agalactiae</i> being the cause of mastitis in cattle besides being an important human pathogen. The observed substrate preference for human IgG1 might reflect that most human invasive <i>S. agalactiae</i> isolates represent distinct subtypes from bovine isolates, as it has been suggested in a temporally and geographically matched isolate characterization study [40]. IgdE<sub>pseudoporcinus</sub> showed double specificity for both human IgG1 and porcine IgG (Fig 2). Since <i>S. pseudoporcinus</i> is closely related to the pig pathogen <i>S. porcinus</i>, the specificity for porcine IgG is not that surprising. It is, however, astonishing that IgdE<sub>pseudoporcinus</sub>, being a human pathogen, has evolved the ability to cleave human IgG1, despite the close relationship to IgdE<sub>porcinus</sub> that does not possess this ability. Thus, it seems advantageous for <i>S. pseudoporcinus</i> to have a human IgG1 degrading protease, underlining the importance of IgG1 in immune responses towards bacterial pathogens. Mutational studies of

**Fig 4. IgdE<sub>agalactiae</sub> and IgdE<sub>pseudoporcinus</sub> are specific for human IgG compared to IgM and IgA.** 0.5 mg/ml human IgG, IgM and IgA were incubated for 16h at 37˚C with (+) or without (-) 20 μg/ml purified rIgdE<sub>agalactiae</sub> (A) or 5% soluble fraction of E. coli cells expressing rIgdE<sub>pseudoporcinus</sub> (B). Reactions were analyzed by SDS-PAGE under reducing conditions. SDS-PAGE was stained with Coomassie blue. Order of lanes within SDS-PAGE was adjusted.

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Fig 5. IgdE_{agalactiae}, IgdE_{pseudoporcinus} and IgdE_{equi} are IgG subtype specific. (A) 0.25 mg/ml human IgG subtypes were incubated for 16h at 37˚C with (+) or without (-) 20 μg/ml purified rIgdE_{agalactiae}. Reactions were analyzed by SDS-PAGE under reducing conditions. IgG cleavage (*) occurred only upon incubation with IgG1. SDS-PAGE was stained with Coomassie blue. Order of lanes within SDS-PAGE was adjusted. (B) 0.25 mg/ml human IgG subtypes were incubated for 16h at 37˚C with (+) or without (-) 5% soluble fraction of \textit{E. coli} cells expressing rIgdE_{pseudoporcinus}. Reactions were analyzed by SDS-PAGE under reducing conditions. IgG cleavage (*) occurred only upon incubation with IgG1. SDS-PAGE was stained with Coomassie blue. (C) 0.09 mg/ml recombinant equine IgG subtypes were incubated for 16h at 37˚C with (+) or without (-) 5% soluble fraction of \textit{E. coli} cells expressing rIgdE_{equi}. Reactions were analyzed by SDS-PAGE under reducing conditions. IgG cleavage (*) occurred only upon incubation with IgG7. SDS-PAGE was stained with Coomassie Fluor Orange Protein Gel Stain. Images of different SDS-PAGE run in parallel have been assembled into one figure.

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Fig 6. IgdE family proteases cleave IgG in the hinge region. (A) The cleavage sites within IgG molecules were determined through N-terminal Edman sequencing of the 32 kDa IgG cleavage products generated by IgdE family proteases. The identified aa sequences (bold) were found in the hinge regions of the respective IgG heavy chains. Homodimer disulfide bond cysteine residues are underlined. 10 aa N- and C-terminal from the identified cleavage site (scissor symbol) of the respective IgG heavy chain are shown. Porcine IgG4a was chosen as a representative for porcine IgG. (B) Sequences of the hinge regions and adjacent parts of the CH1 and CH2 domains of human, porcine and equine IgG subtypes.
IgdE<sub>pseudoporcinus</sub> and IgdE<sub>porcinus</sub> might be able to dissect the residues that mediate this substrate specificity.

We suggest that substrate specificity of IgdE family proteases contribute to the host tropism of some <i>Streptococcus</i> species. Co-evolution of streptococcal opportunistic pathogens and their host could be reflected on a molecular level by co-evolution of IgdE family proteases and their substrate IgG heavy chain molecules. This is highly reminiscent of the co-evolution described between IgA and bacterial proteins targeting IgA, such as IgA-binding proteins and IgA-specific proteases, where reiterative episodes of natural selection are predicted to have shaped the interactions between the IgA and the bacterial proteins, reflecting an 'arms race' [41, 42].

The subtype specificity of the IgdE family proteases of <i>S.</i> <i>agalactiae</i> and <i>S. pseudoporcinus</i> towards human IgG1 and the IgdE family protease of <i>S.</i> <i>equi</i> subsp. <i>zooepidemicus</i> towards equine IgG7 is striking and surprising (Fig 5). The evolutionary benefit for streptococci to possess such IgG subtype specific proteases compared to proteases with broader specificity is at the first glance puzzling. Cleavage of these IgG subtypes might, however, be sufficient to overcome key immune defense mechanisms in certain niches. For example, <i>S. agalactiae</i> is a common cause of invasive neonatal infections in humans [43] and human IgG1 is, along with IgG3, the major human Ig transported across the placental barrier [44]. Thereby cleavage of human IgG1 might be sufficient to evade the Ig mediated immune response in the newborn human host. Moreover, targeted disruption of IgG7 function by <i>S. equi</i> in the horse is likely to significantly comprise IgG-mediated protection, given that IgG7 is one of the predominant subclasses in equine serum [45]. Due to the high diversity in the hinge region of different IgG subtypes [17, 46] it might also be difficult to evolve IgG degrading proteases with broader specificity. Since some IgG subtypes, for example human IgG4, are believed to mediate tolerance [47, 48] it might even be beneficial for an opportunistic pathogen to carry proteases incapable of cleaving these IgG subtypes. Investigations on the potential role of IgdE<sub>agalactiae</sub> in immune evasion in the neonatal and adult host are currently ongoing in our group.

IgdE<sub>equi</sub> is the third IgG degrading protease of <i>S. equi</i> subsp. <i>zooepidemicus</i> described, beside IdeZ1 [5] and IdeZ2 [49]. IgdE<sub>equi</sub> is highly specific for equine IgG7, while both IdeZ1 and IdeZ2 have broader specificity, cleaving IgG of several host species. The abundance of genes encoding IgG degrading proteases in <i>S. equi</i> subsp. <i>zooepidemicus</i> implicates a special importance of an IgG cleaving phenotype of this species. These IgG-degrading proteases might be regulated by different gene regulation systems and thereby expressed during different stages of infection or colonization.

All IgdE family proteases recognize IgG as substrates although the amino acid sequences at the cleavage sites in the respective hinge regions are quite diverse (Fig 6). Therefore preference for IgG as substrates of IgdE family proteases might not only be conferred by the actual cleavage site, but also by motifs lying adjacent to it or within the Fc fragment or F(ab) fragment. Indeed, this possibility mirrors that observed with certain human IgA1 proteases in that residues within the Fc region of IgA have been shown to be essential for recognition of human IgA1 as a substrate for cleavage [50, 51]. Further parallels with IgA1 protease cleavage of IgA1 hinge and the cleavage of IgG hinges by IgdE family proteases described here can be noted. Different IgA1 proteases are known to cleave at different specific peptide bonds in the IgA1 hinge sequence, and evidence suggests that for cleavage to occur each protease has a requirement for the Fab and Fc regions to be separated by a particular number of amino acids, presumably to allow appropriate access and orientation of the protease [52]. Possibly similar spatial
considerations may impact on the ability of IgDE family members to cleave their respective substrates, and may provide a further explanation for their exquisite specificity for particular IgG subtypes.

IgdEagalactiae has the same cleavage site in human IgG1 as papain [53]. However compared to papain, IgdEagalactiae is highly specific for human IgG1 and has only one distinct cleavage site within the heavy chain. Interestingly, this cleavage site is not shared by IgdEpseudoporcinus that instead cleaves the heavy chain two residues closer to the C-terminus, just N-terminal of the putative homodimer disulfide bond cysteine residue. Differential cleavage sites implicate that targeting IgG1 has evolved independently in these two proteases, highlighting the importance for the bacteria to counteract IgG1. The cleavage sites of IgdEporcinus, IgdEpseudoporcinus and IgdEequi have a CpxCP motif just C-terminal of the cleavage site in common. This motif can, however, also be found in many IgG heavy chain molecules that are not substrates of any investigated IgDE family protease, supporting the idea that substrate specificity is determined by features others than cleavage site sequences.

Secreted IgG degrading proteases have been shown to be protective antigens in experimental vaccine and infection studies with S. suis in pigs [54] and S. equi in horses [55]. The described IgDE proteases might therefore be suitable vaccine targets. Given the homology of these proteases, especially in regions close to the active site, vaccination might even give cross protection against several Streptococcus species. This would especially be desirable in the cases of S. agalactiae and S. pseudoporcinus in humans and S. suis and S. porcinus in pigs. Antibodies elicited by such vaccines might both neutralize the proteolytic function of these potential immune evasion factors and potentially mediate antibody dependent cell cytotoxicity against the streptococcal pathogen.

Lastly IgDE family proteases with pronounced species and subtype specificity might also be of biotechnological or therapeutical use, i.e. similar to what has been proven for the IgG degrading enzyme of S. pyogenes IdeS [56–61].

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