The Q rule in Bacteriodetes and the identification and characterization of Porphyromonas gingivalis Glutaminyl Cyclase.

John Andrew Houston
University of Louisville

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THE Q RULE IN BACTERIODETES AND THE IDENTIFICATION AND CHARACTERIZATION OF PORPHYROMONAS GINGIVALIS GLUTAMINYL CYCLASE

By

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B.S. Samford University
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A Dissertation
Submitted to the faculty of the Graduate School
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For the degree of

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THE Q RULE IN BACTERIODETES AND THE IDENTIFICATION AND CHARACTERIZATION OF PORPHYROMONAS GINGIVALIS GLUTAMINYL CYCLASE

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A Dissertation Approved on

November 30th, 2018

By the Dissertation Committee:

Director: Dr. Jan Potempa
Dr. Richard Lamont
Dr. James Graham
Dr. Don Demuth
Dr. David Scott
DEDICATION

This dissertation is dedicated to my parents, Jimmy Allen Houston and Melinda Harris Houston, who have given me unlimited educational potential and the drive to succeed and most importantly my wife, Emily Meadows Houston who has given me endless support and boundless encouragement to continue my passion.
ACKNOWLEDGEMENTS

I would like to especially thank my mentor Dr. Jan Potempa for his unwavering support and endless patience. Without your teaching and assistance none of this body of work could have ever been possible. I would also like to thank the members of my committee: Dr. Richard Lamont, Dr. David Scott, Dr. Don Demuth, and Dr. James Graham for their support and help these past seven years. I would like to also acknowledge the faculty and administration of the University of Louisville School of Dentistry for making this dual degree program a reality and facilitating its integration with my clinical coursework. It has been a journey fraught with peril but yet we emerge relatively unscathed. I would like to thank the many members of Dr. Potempa’s lab, who have taught me so much. Lastly, I would like to acknowledge my wife Emily, who has stood by me through the long nights, early mornings, and stressful days with open arms and warm smiles. This work is for you.
ABSTRACT

THE Q RULE IN BACTERIOODETES AND THE IDENTIFICATION AND CHARACTERIZATION OF PORPHYROMONAS GINGIVALIS GLUTAMINYL CYCLASE

John Andrew Houston

November 30th, 2018

Background: *Porphyromonas gingivalis*, a major pathogen associated with chronic periodontitis, secretes variety of proteins, majority of which begins with glutamine. Several of these proteins were found with pyroglutamate (pGlu) at N-terminus suggesting the presence of this posttranslational modification pathway in *P. gingivalis*. The observation that N-terminal glutamine is over-represented as the first amino acid after signal peptide cleavage, and subsequent confirmation of pGlu formation on the nascent protein via mass spectrometry, led us to conclude that an enzyme must be present as the executor of this reaction. Hypothesis: PG2157 is a glutaminyl cyclase and is responsible for the cyclization of N-terminal glutamine residues. Methods: A homology search was used to identify a gene (PG2157) encoding a protein homologous to human glutaminyl cyclase (QC) in the *P. gingivalis* genome. The gene was cloned, expressed in *E. coli* and recombinant PgQC purified. The protein was crystalized, and structure determined by molecular
replacement. The rPgQC activity was characterized with respect to pH, ionic strength, optimum substrate specificity, and sensitivity to inhibition by an array of non-specific and specific inhibitors. Finally, subcellular localization of PgQC in P. gingivalis was determined. Results: PgQC specificity is restricted for N-terminal glutamine. The enzyme converts this residue to pGlu with $k_{cat}/K_m$ at 1.34 s$^{-1}$. The reaction was fastest at low ionic strength and at pH around 8.0. The activity was inhibited by o-phenanthroline (≥100µM) and EDTA (≥100mM EDTA). Cu$^{2+}$ and Zn$^{2+}$ at ≥100nM exerted ≥90% inhibition. The activity was also significantly affected by cysteamine, imidazole, and reduced glutathione. In bacterial cells PgQC was found associated with the inner membrane as a lipoprotein facing the periplasm. The crystalline structure of PgQC showed strong similarity to human QC on the atomic level. Nevertheless, an inhibitor specific for human QC had a limited effect on the PgQC activity. Conclusions: PgQC is an enzyme resembling mammalian QC and it is responsible for pyroglutamation of proteins secreted by the T9SS of P. gingivalis. This activity is likely essential for bacterium viability since all attempts to produce a viable PgQC knockout failed. Taking into account that also T. forsythia and P. intermedia possess similar enzymes and the frequency of the Q value of Bacteriodetes it is likely that similar post-translational modification plays a pivotal role in protein secretion by these periodontal pathogens. Therefore, inhibition of bacterial QC may represent a novel approach to treat periodontal diseases.
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CHAPTER I: INTRODUCTION

Periodontal Disease

A *periodontal diagnosis* is a term that serves as a crucial determination of a patient’s dental treatment outcome. Clinicians assign these diagnoses on patients as a culmination of the sum of all the clinical information, pertinent medical history, and dental history along with the gross findings from a completed periodontal and oral examination. All the clinical data along with the totality of signs and symptoms are aggregated together and with this sum of information the clinician may arrive at a diagnosis. Additionally, sometimes in more complicated cases, additional lab tests or supplemental information can be instrumental in coming to the correct conclusion. As a clinician, treatment of plaque-induced periodontal diseases generally results in the resolution of the periodontal infection. Also, it is vital to understand that periodontal treatment generally changes the pretreatment diagnosis to a usually more favorable post-treatment diagnosis. To demonstrate, effective prophylactic treatment routinely converts mild, moderate, and severe plaque-induced gingivitis into a state of periodontal health, when stressed with adequate home care instructions. To further illustrate this point, successful treatment of plaque-induced periodontitis, when followed up at re-evaluation appointment, will often be converted to a state of periodontal health with reduced periodontium, indicating that treatment has achieved the goal of creating shallower pocket depth measurements.
Periodontal disease is term that encompasses multiple differential modalities into one general disease term. Within the diagnosis of periodontal disease, there are more diverse classification systems that both clinicians and researchers alike use. The most descriptive and thorough clinical classification system was described by Armitage in 1999, and is still utilized by clinicians today [1].
Table 1 Presentation of Periodontal Diseases in Humans

<table>
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Armitage [1-3]
Most patients who present with plaque-induced periodontitis will in fact possess the chronic form of the disease [4-7]. The most likely presentation of a chronic periodontitis patient is a patient greater than 30 years of age; gross plaque and calculus, substantial gingival inflammation is present, deepened periodontal pockets (> 3mm), and the presence of calculated periodontal attachment loss. Generally speaking chronic periodontal disease is a slow continuously progressing disease [8-10], but within this process may exist short periods of advancing rapid attachment loss [11, 12]. Previously chronic periodontitis was given the misnomer “adult periodontitis” since only the adult population was believed to be susceptible to the disease. However, this has been shown to be false as a result of past and more recent epidemiologic data clearly showing that chronic periodontal disease can be present in younger populations [9, 13]. Although specifically chronic periodontitis can be observed to occur in either localized or generalized distributions within patients, these specific two forms appear to be vastly similar or almost identical with regards to their presentation, pathogenesis, and progression. Aggressive periodontitis, however, is far more rare than chronic periodontitis and mainly affects children and adolescents. It also can occur in both localized and generalized presentations, but unlike chronic periodontitis, these distributions differ significantly with respect to their pathogenesis [4, 13]. Localized aggressive periodontitis (LAP) and generalized aggressive periodontitis (GAP) were previously called “localized and generalized juvenile periodontitis”, respectively [14].
Biofilms

These different presentations of periodontal diseases are best described as multifactorial infections, which are elicited by the presence of an entire community of complex of bacterial species. This community of organisms referred to as a “biofilm” interacts with the host, specifically with connective tissue and immune cells, which in turn causes a release of large number of pro-inflammatory cytokines and chemokines. These pro-inflammatory mediators lead to the degradation and destruction of cellular structures of the periodontal organ, namely: bone, periodontal ligament, sulcular epithelium, and junction epithelium [15]. The trigger for the initiation of periodontal disease is the presence of dysbiotic microbial biofilms composed of bacteria that colonize the tooth surface in the sulcular region. This region lies between the tooth surface below the cement-enamel junction and the gingival margin and plaque formation is mediated through specific adherence interactions and accumulation due to architectural changes in the sulcus [16]. The characteristics of microbiological progression from periodontal health to gingivitis (e.g. chronic inflammation of the gingival tissue without tissue destruction), and eventually to periodontal disease are vast and complicated [17].

Although it has been previously estimated that out of nearly 1,000 different microbial taxa are able to colonize, at least transiently the oral cavity of humans and from a complex organized biofilm [18], it is still not fully understood how the
multitude of different species exist and work harmoniously to begin the pathogenic process and progress to full blown disease state. Previous studies have shown that of the just under 1000 species detected in the oral cavity approximately 500 are found to be present in periodontal plaque [19]. Some of these organisms most certainly are simply commensal or opportunistic species, but it is well documented that certain species drive the pathogenicity of the bacterial plaque into more virulent and destructive states [20]. Previous work has shown that even early colonizers and previously deemed commensals such as *Streptococcus gordonii* can have a profound effect on the pathogenicity of other organisms such as *A. actinomycetemcomitans* and *Porphyromonas gingivalis* [21]. It has been shown in several studies, and mentioned previously here that a multitude of bacterial species call the human mouth their home, humans overall understand very little about the consequences of harboring our bacterial inhabitants [22, 23]. Can the presence of just one of the “bad guys” turn our whole “neighborhood” into a difficult place to manage? Or does it take a combined effort of the community as a whole to corrupt our own host defenses. With regard to other dental diseases, such as dental caries, evidence is strong enough to allow us to point to the causative agent; *Streptococcus mutans* and *Lactobacillus sp*. Since these species have been identified, many papers have assessed the bacterial causative agents in patients in the varying states of periodontal health, gingivitis, and periodontitis [17, 20, 22, 24, 25]. We are aware of a great many gram-positive bacterial species that serves as beneficial commensal species and serve to help maintain important oral health. We now also know that as the progression of the disease proceeds and heads
further towards gingivitis and chronic periodontitis, greater numbers of gram-negative species begin to inhabit the sulcular epithelium and surrounding dentoalveolar structures. More striking though, is the simple fact that there is still no verifiable shown data, that one solitary species can solely cause the wide-ranging effects that periodontal disease can elicit in the mouth. No one organism can account for the varied destructive processes that occur in the disease process. This leaves us with the thought that the disease can only progress if several bacterial species join forces and the signs and symptoms of periodontal disease are in fact the result of a group effort. It must be clear then that a specific “consort” or “complex” of bacterial species must trigger the transition from a state of oral health to periodontal disease. Innumerable studies have focused on one specific bacterial species responsible for signs and symptoms of periodontal disease, but these do not account for the whole story. This, taken as a whole, supports the notion that the effects from periodontal disease must be the result of consortia of bacterial species acting as a complex biofilm to cause, elicit, and promote the disease [24-27]. Recently there has been reassessment of the “roles” of several bacterial species that are known to be involved in the progression of periodontal disease. These organisms were routinely found in examinations of patients who presented with periodontal disease and also in healthy controls. These species are now even commonly referred to as periodontopathogens. Included in this group are *P. gingivalis, A. actinomycetemcomitans, T. forsythia, and T. denticola*. Influential studies by Socransky and Haffajee [24] used newer methodology of stratifying these bacteria into their respective niches based on their roles within
oral microbiomes in the disease process leading to change from health to disease states. Researchers organized the organisms into groups or “complexes”. Individual complexes were based on the prevalence of the bacterial “affiliations” with one-another, and corresponding complexes associations with health, gingivitis, or periodontitis disease conditions [25-29]. These groups or complexes of different microbes were also stratified according to the sequence of colonization on the tooth surface in conjunction with periodontal disease severity. What Socransky et al. [20] labeled as the ‘red complex’ contained bacterial species that show up later in biofilm maturation phase. These bacterial species, namely, \( P. \) gingivalis, \( T. \) denticola, and \( T. \) forsythia (previous names \textit{Bacteroides forsythus} or \textit{Tannerella forsythensis}) were shown to be effective periodontopathogens [11, 20, 24, 26, 30]. In this same train of thought these investigators also concluded that this “red complex” represents the pinnacle of biofilm maturation and development and thus leading to advancing disease states.

Multiple other researchers have made note of this cooperation between species within the same complex and between members of different complexes. A strong association has been observed between \( T. \) forsythia and \( P. \) gingivalis found in periodontal pockets in patients, whilst in fact \( P. \) gingivalis has not been detected in the absence of \( T. \) forsythia within the periodontal pocket [31]. A strong relationship between \( P. \) gingivalis and \( T. \) denticola was found whilst taking plaque samples gathered in a study looking at diverse ethnic groups [32]. Moreover, constituent members of the red complex were located in significantly higher percentages in patients with periodontitis [33, 34] and also in probing sites with deeper pockets.
These separate bodies of work support the theory that no one single bacterial species is etiologic for periodontal disease propagation or progression, and periodontal disease is not a singular “bacterial infection” but instead that the sum total of bacterial species in the oral cavity co-existing and co-habituating the oral microbiome are required to initiate the onset of periodontal disease. How these different species and organisms, as well as virulence determinants of the individual bacteria, contribute to disease progression remains unclear. Another hypothesis in more recent years has challenged the way we typically look at the “complexes” of oral microbiota in the mouth. While looking at the red complex organisms one would beg to ask the question, would higher numbers of these organisms correlate with greater periodontal tissue destruction. In fact, certain pathogens, more specifically, some from the red complex are actually found to be present in low numbers. More recently, multiple analyses of the human microbiome have started to shift the focus on biofilms away from consideration as a true “infection” but instead as a dysbiotic disease. We are beginning to live in the age of the dysbiotic inflammatory disease. Dysbiosis is a process by which communities of normally healthy bacterial species become unstable and unregulated and in response drive the inflammatory process. This outlook has become more optimal a term to assign to periodontal disease than just giving few species the honor of being disease causing agents. In healthy individuals, oral microbiota remains in a state of harmony and regulation, but in susceptible individuals or compromised patients, dysbiosis can lead to unfavorable and harmful host-microbial interactions and eventually leads to periodontal inflammation and destruction of the periodontal
tooth supporting structures [41-43]. This in turn leads to the inflammatory reaction products being released into the gingival crevicular fluid (CGF) which then promulgates and promotes furthering the disease process [44-46]. This cycle of inflammation and dysbiosis could be a key factor that plays into periodontal disease’s main components, its lasting effects and difficulty to treat. Traditional therapy of scaling and root planning for periodontal disease, along with debridement and maintenance of the oral cavity may now be possibly augmented to include adjuvant therapies that target this dysbiotic shift towards more pathobiont species [47, 48]. In addition to this potential benefit, some of released host factors or molecules could have utility as disease markers in patient treatment. This could serve to reflect a more accurate disease state or give a better indication of the level of periodontal inflammation and disease progression than was previously available by solely clinical examination [49].

The keystone pathogen hypothesis was posited as a result of new information coming to light in the field. Studies showed that conversely, relatively low level of the red complex species were actually present in biofilms, and in the mice bone loss model suspected periodontopathogens were only needed in small numbers to influence periodontal inflammation and bone loss [50, 51]. The small bacteria cell count were enough to initiate a shift in the number and makeup of the complex biofilm. This alteration of the biofilm composition occurred before the onset of significant bone loss and was closely associated with the colonization of \( P. \) gingivalis, indicating that the cause of the disease was the shift in bacterial species
composition or dysbiosis not solely the offending organisms' presence alone [51]. This in conjunction with the observed fact that *P. gingivalis*, in the absence of other commensal bacteria, fails to cause periodontitis in mice. For the purposes of this dissertation, our discussion will focus mainly on the periodontopathogen and keystone pathogen, *P. gingivalis*, its associated virulence factors, and the methods by which these are manufactured and undergo post-translational modifications.

*Porphyromonas gingivalis*

*P. gingivalis*, as mentioned above, has been a heavily studied pathogen in the oral cavity and will be mainly the subject of this dissertation. As mentioned previously we label this bacterium as a keystone pathogen in human periodontitis [50, 52]. By placing this moniker on this bacterium, we imply that this organism is capable of causing dysbiosis. The dysbiosis can manifest in the form of either relative number or abundance of pathogenic species or change in the role or stratum of the species within the biofilm, even at low microbial levels within the biofilm. Microbial analysis studies have shown that despite its known importance in oral biofilm, *P. gingivalis* is only a minor constituent of periodontal disease-associated biofilms [53-55]. This is further confirmed by studies showing that, in a mouse model of periodontitis, *P. gingivalis* was able to colonize in low-levels and was shown to cause an increase in certain populations of the periodontal microbiota followed by inflammation-driven alveolar bone loss [51]. In addition, serving as a keystone pathogen comes with certain responsibilities such as serving a specialized role in the biofilm community. This role coordinates and modulates the activity of other organisms within the plaque and plays an essential role in its pathogenicity [56, 57]. In this
role, \textit{P. gingivalis} influences the transition from mainly commensal bacteria into a pathogenic biofilm [52]. \textit{P. gingivalis} is a Gram-negative, obligate anaerobe, and asaccharolytic rod which possess a number of virulence factors [58-60]. Many of these virulence factors are directly related to \textit{P. gingivalis’} ability to subvert the host immune system. The use of these virulence factors is what makes \textit{P. gingivalis} an effective pathogen.

\textbf{Gingipains}

Some of these virulence factors alluded to previously are the proteases, gingipains. Included in this category are the arginine-specific gingipains [Arg- gingipain-A (RgpA) and Arg-gingipain-B (RgpB)] and also the lysine-specific gingipain [Lys- gingipain (Kgp)]. These are encoded respectively by their three constituent genes within the \textit{P. gingivalis} chromosome commonly referred here as \textit{rgpA}, \textit{rgpB} and \textit{kgp} which are heavily conserved among different clinical and experimental strains of \textit{P. gingivalis} [61]. The products of the translation of these genes (specifically the \textit{rgpA} and \textit{rgpB}), RgpA and RgpB, both contain a caspase-like domain (that retains specificity for Arg-Xaa peptide bonds) and an immunoglobulin-like domain. Uniquely in the proteinase RgpA, the protease and Ig-like domain is subsequently followed-up with a large hemagglutinin-adhesin C-terminal extension. Closely related to the above protein, the \textit{kgp} gene-translation product, Kgp, contains a catalytic domain specific for Lys-Xaa peptide bonds and also contains a hemagglutinin-adhesion domain not too unlike the \textit{rgpA} translation product. [62, 63], RgpB lacks the hemagglutinin-adhesin domains but the short C-terminal domain is conserved. Gingipain translation products undergo heavy post-
translational modification. Nascent translation products encompass a pro-
fragment, a catalytic domain, an Ig-like domain, hemagglutinin-adhesion domains
(only in RgpA and Kgp) and the conserved C-terminal domain (CTD). During
secretion of RgpB, the pro-fragment is proteolytically removed and subsequent
processing of the CTD reveals the active catalytic domain followed by the Ig-like
domain. Subsequent glycosylation and incorporation to the outer membrane
surface occurs afterwards. In the case of RgpA and Kgp similar processing occurs
but the hemagluttanin-adhesin domains remain non-covalently associated with the
catalytic domain. Lastly, gingipains once processed are secreted either as a
monomeric form specific to the case of RgpB, or as complexes of protease and
hemagglutinin-adhesin domains in the case of RgpA and Kgp. These complexes
are either predominantly attached to the bacterial surface or released into the
medium in a soluble form. This is dependent on a P. gingivalis strain, for example
the strain HG66 secretes soluble gingipains freely into the media. Gingipain
activities serve as meditators for nutrient acquisition, serve to cleave receptors on
the host cell surface, and moreover, avoid and subvert the host immune system
by inactivation of cytokines and components of the complement system. One of
the most profound changes observed in patients with clinical periodontal disease
is the aberrant remodeling of the host periodontal tissues. Structural changes that
are often associated and observed with advanced periodontal disease include
alveolar bone resorption and periodontal ligament destruction. These changes in
structure in turn lead to attachment loss and periodontal pocket formation and
eventually tooth loss. Periodontal pockets found in patients with periodontal
disease are also lined with epithelial cells that have had alterations, making them distinctly different from healthy cells that form the junctional epithelium found in the healthy periodontium. To accomplish this tissue-remodeling host cells require proteolytic degradation of important structural elements and extracellular matrices such as the collagen fibers forming the periodontal ligament and proteins involved in cellular junctions and extracellular matrix proteins [64-67]. The mechanism by which gingipains play a role in this process is complex and beyond the scope of this introduction. But in summary gingipains are important both directly and indirectly involved in the pathological tissue remodeling associated progressing disease process of periodontitis. Nevertheless, it is most likely that gingipains are not solely involved as the major virulence factor in periodontal disease tissue destruction [59]. It is instead much more probably that these proteases synergize with other mechanisms and virulence factors used by *P. gingivalis* and other bacteria and disrupt the host proteolytic balance and interfere with endogenous host protease inhibitors. Once this balance has been disrupted, host proteases, namely MMPs, TIMPs, and neutrophil elastase, drive the response in the periodontium for accelerated remodeling and pathological destruction of the host tissues[64, 68]. The effects of gingipains have been studied in great details and the role that they play in the disease process is expanding [59]. This variable output of potent virulence factors gives *P. gingivalis* quite an arsenal with which to exert effects into the host organism and biofilm community.

**Protein Secretion Systems of Bacteroidetes Species**
Apart from the Sec pathway that exports proteins into or through the inner membrane [69, 70], the armamentarium of well-characterized secretion pathways of Bacteroidetes is surprisingly limited. Bacteroidetes possess both type 1 and type 6 secretion systems (T1SS and T6SS) that are responsible for secreting proteins to extracellular environment bypassing the periplasm. Recently, there has been a great deal of effort produced to characterize the T9SS. This system is typical for Bacteroidetes and rarely, if ever, found outside the phylum [71, 72]. The T9SS machinery transports proteins across the outer membrane, which were first exported into the periplasm via the Sec translocon. All proteins secreted by T9SS possess a characteristic C-terminal domain (CTD) that codes for transportation through the outer membrane [71, 73]. The processes that are undertaken upstream of CTD-dependent secretion have garnered less investigation, mainly due to their ubiquity in other organisms and Bacteroidetes and have been extensively studied in gram-negative bacteria in general [70]. We report here that these upstream processes are unique to Bacteroidetes and do in fact merit attention. To put this more into perspective, some background on the Sec translocon is helpful.

The Sec pathway identifies target proteins by the presence of a signal peptide [74]. These signal peptides contain a tripartite architecture. They consist of a positively charged N-terminal region, which is thought to designate proteins to the phospholipid membrane, a hydrophobic region, which is hypothesized to be inserted into the membrane, and finally a shorter region that often contains a
consensus motif for a signal peptidase [75, 76]. Proteins that contain signal peptides can escape from the Sec translocase in two different ways. If they manage to escape “laterally”, they will then become contained within the inner membrane [69]. The other way would have them reach the periplasm, but the signal peptide portion is not cleaved and remains in the inner membrane. This leaves the proteins’ C-terminal end (with a signal peptide cleavage site) exposed on the periplasmic surface of the inner membrane [75]. The eventual fate of proteins that reach this stage will depend on the type of the signal peptide contained within.

Proteins that carry a type I signal peptide are released from their membrane anchored signal peptide by signal peptidase I (SPI) [74, 75]. They will subsequently remain in the periplasm, or if necessary be transported further, for example, by a T9SS through the outer membrane, ultimately destined for the surface of the outer membrane or for release into the environment [71]. Proteins which carry a type II signal peptide undergo processing differently. A diacylglycerol transferase (termed Lgt) first attaches a diacylglycerol membrane anchor to the cysteine residue which resides immediately downstream of the signal peptide [77]. Secondly, signal peptidase II (SPII) (also known as lipoprotein signal peptidase or Lsp) can cleave a lipoprotein upstream of the modified cysteine residue [78, 79]. Occasionally, lipoproteins can remain attached to the inner membrane while others are transported to the outer membrane via the Lol system [80]. Thus, lipoproteins
found in gram-negative bacteria are generally periplasmic proteins which are anchored either to the inner or outer membrane of the bacteria.

The N-terminal residue of proteins is frequently chemically modified, and these modifications often have a signaling role. If an N-terminal glutamine residue is exposed as a result of proteolysis, these glutamine residues have the capability to spontaneously cyclize to pyroglutamate, with concomitant release of ammonia as a side product. The reaction is also facilitated by inorganic catalysts such as phosphate ions serving as the proton shuttle, and furthermore, can be catalyzed enzymatically by glutaminyl cyclases (QCs) [81].

**Glutaminyl Cyclase**

As one of the enzymes for protein post-translational modifications, glutaminyl cyclase (QC; glutaminyl-peptide cyclotransferase (QPCT), EC 2.3.2.5) is an acyltransferase that catalyzes N-terminal pyroglutamate (pGlu) formation on proteins or peptides and the concomitant release of ammonia or water molecules see Figure 1 [82].
Fig. 1: N-terminal cyclization of glutaminyl peptides by QC. Glutaminyl containing peptide is cyclized in the presence of QC by formation of an intermediate state that culminates in the release of ammonia.
The cyclization of L-glutamine into pyroglutamate is considered a quasi-
irreversible intramolecular acyl transfer reaction. The N-terminal peptide amino
group is responsible for a nucleophilic attack on the γ-carbon amide, forming a
tetrahedral intermediate. Subsequent decomposition of this intermediate occurs
from net 1,3-proton transfer from Nα onto a leaving ammonia, forming the pGlu
ring. This cyclization or modification appears to be involved in the structural
stability of secreted proteins, resistance to aminopeptidase degradation, and
hypothetically play an important role in mediating receptor binding [83]. QC is a
catalytically non-discriminatory enzyme; acting upon an available N-terminal
 glutamine residue, although it does exhibit slight preference for substrates
containing a penultimate bulky hydrophobic residue [84]. Recently it has been
shown that phosphate ions in conjunction with QC are able to operate as a
coordinated proton transfer system. This system is able to effectively increase the
rate of cyclization by QCs. Possibly; phosphate acts as a dual functioning acid–
base catalyst to increase the proton transfer rate, where simultaneous protonation
and deprotonation occur together in conjunction with a cyclic-ring transition state
(Fig. 2 A and B). Previously phosphate ions have been hypothesized to serve as
dual functioning catalysts during the iminolactone hydrolysis [85] or for thiamin
enzyme tautomerization [86]. Acid–base catalysis is the primary source of
catalysis used by QCs. The proposed role of the active-site zinc ion in QC can be
nicely correlated to the above-described rate-limiting trapping of $T^\pm$ in acyl
transfer reactions in solution (Fig. 2 B). The active site zinc functions as a strong
Lewis acid, and proton transfer from the attacking α-amino group to the leaving
ammonia is accelerated by a glutamic acid moiety in conjunction with two other acidic residues, as suggested by QC crystal structures [87, 88].
Fig. 2: Hypothetical role of phosphate (A and B) and QC (C) in acceleration of glutaminyl cyclization. Phosphate ions facilitate the proton transfer onto the leaving ammonia. The catalysis might involve facilitated proton transfer onto the oxanion (A) or deprotonation of the attacking α-amino group (B). The proton transfer in QC is facilitated by a glutamic acid residue. A significant acceleration of the reaction is achieved by “trapping” of the oxanion by the active site zinc ion. Possibly, the proton transfer from the attacking alpha-amino group to the leaving ammonia occurs simultaneously [89].

QC was originally discovered in dried latex belonging to the *Carica papaya* (CpQC) plant species [90]. Glutaminyl cyclases (QCs) have been demonstrated in multiple plant and bacterial species and share a common ancestry [91-94]. It has been
found that the known mammalian QCs are either Golgi-resident or freely secreted enzymes [93]. Several peptide hormones and proteins carry N-terminal pyroglutamyl residues. Until somewhat recently, the cyclization of L-glutamine peptides’ N-terminus into pyroglutamate was thought to occur spontaneously [95]. This cyclization was shown that the conversion occurs under physiological conditions but at a very slow rate, therefore necessitating the need for enzyme catalysis [96]. Mammalian and plant QCs, however, have been discovered to serve as this executor of the reaction [97-101]. In general, both mammalian and plant QCs appear to have similar molecular masses, ~33 and ~40 kDa, respectively, and both are monomeric proteins [102, 103]. The primary structures of the proteins, however, display no sequence similarity, and the secondary structure of the individual QCs are completely different. The plant CpQC and the related QC from bacterial plant pathogen *Xanthomonas campestris* (XcQC) adopt a five-bladed beta-propeller fold [104]. As one would expect for beta-propeller proteins, the active site is located near the propeller axis [105]. Mammalian QCs however are shown to encompass an α/β-fold [106-108]. In addition, currently identified plant QCs do not share sequence homology to other known plant enzymes, which places them into a separate enzyme family or subfamily [100]. On the other hand, identified mammalian QCs possess clear homology toward known bacterial aminopeptidases, thus suggesting possible the evolutionary origin of the protein family [107]. The mechanism of catalysis for plant QCs is much less clear than for mammalian QCs. In mammalian QCs, the cyclization reaction serves its function during maturation of numerous cytokines and neuropeptides in the secretory
pathway, such as gonadotropin-releasing hormone (GnRH), thyrotropin-releasing hormone (TRH), and monocyte chemotactic protein-2 (MCP-2). The theoretical role of the pGlu residue on these peptides is believed to serve two functions; (1) protecting the peptides from peptidase degradation and (2) providing the proper conformation of the peptide to facilitate receptor binding [109, 110]. Uncontrolled expression of QC in humans (HsQC) has been shown to be related to certain pathological conditions, for example Alzheimer disease [111]. Site-directed mutagenesis and X-ray crystallography studies of recombinant HsQC have given us large amounts of information about the catalytic mechanism of QCs [112, 113]. Furthermore, HsQC has been used as a target for inhibitors, which have been synthesized and developed as drugs to treat the relevant diseases that occur from their overabundance [94, 114]. QCs have also been shown to be present in a number of snake venom isolates [115] in keeping with finding that proteins and toxins in snake venoms are often resistant to Edman-degradation during sequence analysis. Cases that have been shown include bradykinin-potentiating peptides [116, 117], metalloproteinase inhibitors [118], and endogenous metalloproteinases [119]. Based on current literature, human QC is thought to share the scaffold of known bacterial aminopeptidases [120]. This is in contrast to the fact that all the putative QCs thus far identified in bacteria share homology with plant QC. When analyzed by atomic absorption spectroscopy, HsQC has been shown to contain one zinc ion per HsQC molecule [121]. Competitive inhibitors of HsQC have been identified, such as heterocyclic compounds (imidazole, tetrazole, and triazole rings). This is most likely due to nitrogen atoms serving as good coordinators of
the active-site zinc ion found in QC [122, 123]. Solely the fact that HsQC could be inhibited/inactivated by heterocyclic inhibitors suggests that the zinc ion located within the QC is essential for catalytic activity [94]. In stark contrast, the same heterocyclic chelators display no inhibition of plant QCs [103]. Considering this information together, both plant and mammalian QCs are thought to belong to two different enzyme families with different ancestral origins [100]. While mammalian QCs are implicated in the maturation of numerous neuropeptides and cytokines, such as thyrotropin-releasing hormone (TRH) and gonadotropin-releasing hormone (GnRH) [102, 124, 125], the physiological function of plant QCs remains poorly characterized. Recently, it has been suggested that plant QCs could be involved in defense mechanisms [126, 127], a hypothesis that is supported by the observation that the quantity of plant QC expressed increases greatly as a result of repeated injury [127]. The role of QCs in both bacteria and parasites has not been studied in depth. Advances in methodology for QC activity assays [128] and the discovery of new HsQC inhibitors [114, 129] have rendered better tools to explore QCs. A new QC has been isolated from *P. gingivalis* and its role has been examined for the first time. Structural homology models of known QCs were generated and compared. Furthermore, we successfully cloned and sequenced the QC cDNAs from *P. gingivalis* and compared the sequence with that of human and other animal QCs. In addition, we also examined the optimal pH range, stability, and the effects of metal chelators, metal ions, and finally examined specific QC inhibitors on the representative bacterial QC. These QCs share similar catalytic activity but structurally remain distinctly different. [113, 126]
In studying the proteome of *P. gingivalis* and its novel T9SS, protein sequences destined for secretion via the T9SS with the conserved C-terminal domains (CTDs) that facilitate secretion from through the outer membrane complex [73]. Approximately 30 CTD-bearing proteins exist in *P. gingivalis* genome and these proteins all contain this conserved C-terminal domain [73]. Recently it has been demonstrated that the CTD-containing proteins are secreted and attached to the cell surface via the type IX secretion system (T9SS) [73, 130-132]. These CTD-bearing proteins undergo extensive post-translational modifications before attachment to the surface. Belonging to this group of CTD-containing proteins are the gingipains (RgpB, HRgpA, Kgp), which are major virulence factors of this periodontopathogen [133]. Upon examination of the sequences of these T9SS cargo proteins it can be observed that a N-terminal Gln residue is present just after the N-terminal signal peptide. Mass spectrometry analysis of these secreted proteins revealed that after post-translational modification, a pGlu residue is present at the N-terminus of the mature protein. This led us to hypothesize the existence of a bacterial QC present in *P. gingivalis*, responsible for this cyclization of Gln to pGlu. The presence of proteins containing N-terminal pyroglutamate residues has been noted in previous proteomic studies on *Bacteroidetes* species [73, 134-137]. At the outset of this study, we became aware of a drastic overrepresentation of glutamine residues after SPI cleavage sites in *P. gingivalis*. Starting from these two observations, we aimed to (1) clarify the pathway of pyroglutamate formation, (2) determine the fraction of SPI substrates that could be cyclized, (3) estimate the fraction that is actually cyclized, (4) test a possible role
of pyroglutamate formation in sorting, initially for \textit{P. gingivalis} only, and then for \textit{Bacteroidetes} in general, (5) characterize the function and importance of PgQC within the T9SS. The function and purpose of the PgQC is hypothesized. In this dissertation, we attempt to verify the existence, explain the purpose and role of QC in \textit{P. gingivalis}, and reveal its potential importance in the secretion pathway in \textit{P. gingivalis} and other \textit{Bacteriodetes}. 
CHAPTER II: MATERIAL AND METHODS

Proteomes and taxonomy information were taken from UNIPROT [138]. Signal peptides and cleavage sites were predicted using the batch version of SIGNALP4.1 [139]. Lipoproteins were predicted using LipoP1.0 [140]. Sequence logos were generated using the program Weblogo [141]. Intersections between proteins with predicted signal peptide and predicted lipoproteins were using the UNIX comm tool. Since not all UNIPROT species have been fully identified, some species names such as “Tannerella sp.” were encountered.

Cloning, expression and purification of recombinant PgQC

The QC protein was expressed as a GST-tag fusion protein. Briefly, the entire coding region of qc identified by BLAST search (PG_2157) was amplified from a P. gingivalis W83 genome template with Platinum Taq DNA Polymerase High Fidelity (Invitrogen) using primers F1_QC: ATTAGAATT CATGAAAAAGACTGATAAAACAGGAG and R2_QC: ATTACTCGAGTCAGTGTGAAGCGGCTTTCACCTGTTCG. The 1001 bp PCR product was digested with EcoRI – Xhol and cloned downstream in-frame with the sequence encoding gluthatione S-transferase (GST), into EcoRI – Xhol digested pGEX-6P-1 expression vector (GE Healthcare). Following confirmation by PCR, resulting expression pGEX/QC vector was transformed into E. coli BI21 (DE3) expression host. Transformed E. coli cells were grown in LB media at 37° C until
OD\textsubscript{600} 0.6, cooled down to 24 °C and expression of recombinant protein was induced with 0.1 mM isopropyl-1-thio-β-galactopyranoside (IPTG). After overnight cultivation, cells were harvested by centrifugation (6,000 x g, 20 min), resuspended in PBS supplemented with lysozyme, and lysed by sonication (3 cycles of 10 x 3 s pulses at 17 W). Cell lysate was clarified by centrifugation (30,000 x g, 30 min) and loaded onto a pre-equilibrated glutathione-Sepharose™ High Performance column. Recombinant GST-QC fusion protein was eluted using 50 mM Tris-HCl, pH 8.0, supplemented with 10 mM reduced glutathione. The purified GST-QC protein was subsequently incubated with PreScission™ Protease (GE Healthcare) and subjected again to chromatography on glutathione-Sepharose™ to remove the GST tag. The purity of the resulting protein was verified by SDS-PAGE electrophoresis (NuPAGE® 4-12% Bis-Tris Gel, Invitrogen). Protein concentration was determined by BCA Assay (Sigma).

**PgQC activity assay**

The activity of PgQC was determined essentially as previously described (33). Briefly, 150 μl of the assay buffer (40 mM Tris-HCl, 400 mM KCl, pH 8.0), 10 μl of chromogenic substrate (200 mM H-Gln-AMC in DMSO), and 10 μl of a recombinant bacterial pyroglutamyl aminopeptidase (25 U/ml, Unizyme Laboratories, Hørsholm, Denmark) were mixed together in a microtitration plate and preincubated 10 min at 30°C. The reaction was initiated by addition of 30 μl appropriately diluted purified rPgQC or *P. gingivalis* whole culture or washed
bacterial cells or subcellular fractions and after 1 min incubation the increase in fluorescence ($\lambda_{ex} = 380$ nm, $\lambda_{em} = 460$ nm) was recorded for 10-20 min at 30°C. Unspecific cleavage of the H-Gln-AMC substrate was determined by omitting pyroglutamyl aminopeptidase, the auxiliary enzyme. If necessary, the unspecific cleavage was subtracted from PgQC activity. Metal-ion inhibition reactions were carried out in similar fashion with each respective inhibitor added to the reaction mixture prior to initiation of the assay.

**P. gingivalis culture and cell fractionation procedures**

*P. gingivalis* culture fractionation was performed at 4°C as described previously [142] starting from stationary-phase (a 2-day-old) cultures adjusted to an OD$_{600}$ of 1.5. Briefly, cells were collected by centrifugation at 6,000 × g for 15 min, washed once with phosphate-buffered saline (PBS), and resuspended in 5 ml of 0.25 M sucrose and 30 mM Tris, pH 7.6. After mixing gently for 10 min cells were repelleted at 12,500 × g for 15 min. The outer membrane was disrupted by the rapid addition of ice-cold distilled H$_2$O and the spheroplasts were pelleted by centrifugation at 12,500 × g for 15 min. The supernatant was designated the periplasmic sample. The remaining spheroplast pellet was resuspended in 5 ml PBS and ultrasonicated in an ice-water bath. Cellular debris and membranes were pelleted by ultracentrifugation at 150,000 × g for 1 h, and the supernatant was designated the cytoplasmic sample. The remaining pellet was washed and resuspended in cold PBS by sonication. This fraction was designated the
membrane sample. For individual membranes separation washed collected cells were lysed by ultrasonication as described above. The membranes were pelleted by ultracentrifugation (150,000 × g, 1 h) washed with PBS to remove periplasmic and cytoplasmic proteins and resuspended in PBS by sonication. The inner membrane was dissolved with Sarkosyl (lauryl sarcosine) and the residual Sarkosyl-resistant outer membranes (OM) were pelleted by ultracentrifugation (150,000 × g, 1 h). The supernatants were designated the IM samples while pellets washed and suspended by sonication in PBS were designated the OM samples. Purity of the various fractions was checked by Western blotting for A-LPS or gingipains and the biotin containing 15 kDa biotin carboxyl carrier protein (AccB alias MmdC or PG1609) as OM and IM specific markers, respectively [62, 143] (data not shown).

**Generation of *P. gingivalis* ΔRgpB deletion mutant**

For subsequent analysis, generation of plasmids suitable for *rgpB* gene mutagenesis the pRgpBall master plasmid was first engineered based on the pURgpB-E construct [142]. A partial *rgpB* gene section upstream of the erythromycin cassette was replaced with whole *rgpB* coding sequence, together with an 817 bp fragment containing its potential promotor. The new fragment was amplified with primers RgpBall_F and RgpBall_R using genomic DNA of *P. gingivalis* W83 and ligated into the linearized pURgpB-E plasmid (with EcoRI and
SmaI restriction enzymes) by the Gibson’s method [144] resulting in pRgpBall-erm plasmid. All primer sequences are listed in Table 2.

<table>
<thead>
<tr>
<th><strong>pRgpBall-erm plasmid</strong></th>
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<tbody>
<tr>
<td>RgpBall_F</td>
<td>GACGGCCAGTGAAATTCTTAAACCAT GCTGTGGTGACGAG</td>
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<tr>
<td>RgpBall_R</td>
<td>AGCGGAAGCTATCCCAACAGTCT CTTGGCGTAGTGCCAA</td>
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<th><strong>pRgpBdel-erm plasmid</strong></th>
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<tr>
<td>RgpBQ1N_Rs</td>
<td>AACGCCATTCTCCCAACAG</td>
</tr>
<tr>
<td>RgpBdelFs</td>
<td>GGGATAGCTCCGCTATTGCT</td>
</tr>
<tr>
<td>RgpBdelRt</td>
<td>GGGTCTGCGGGCTGTGCAACGC CATTCCCTCCCAACAG</td>
</tr>
<tr>
<td>RgpBdelFt</td>
<td>TGCACAGGCCGAGACCCCGGGAT AGCCTCCGCTATTGCT</td>
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<th><strong>RgpBQ24N mutation</strong></th>
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<tr>
<td>RgpBQ1N_Fs</td>
<td>CGGTCGCAACCAACAAATGAC</td>
</tr>
<tr>
<td>RgpBQ1N_Ft</td>
<td>TGCAAACCGCAGACCGCCGTC GCAACCCAACAGTAC</td>
</tr>
<tr>
<td>RgpBQ1N_Rs</td>
<td>AACGCCATTCTCCCAACAG</td>
</tr>
<tr>
<td>RgpBQ1N_Rt</td>
<td>CGCTCTGCGGGTTTGCAACACG CATTCCTCCCAACAG</td>
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<tr>
<th><strong>RgpBQ25A mutation</strong></th>
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<tr>
<td>Q25AF</td>
<td>GGAATGGCGTTTGCAAGCTCCGCAGAGCGCGGTC</td>
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<tr>
<td>Q25AR</td>
<td>GACCCCGCTCTGGCGAGCTGCA ACGCCATTCC</td>
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<th><strong>pNRgpA-tet plasmid</strong></th>
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<tr>
<td>RgA_Up_F</td>
<td>GCCAGTGATATTGTCAGAGAGC CGA</td>
</tr>
<tr>
<td>RgA_Up_R</td>
<td>CGTTGTGGATCTCAGGCTACCAT ATCTTTAACC</td>
</tr>
<tr>
<td>RgpA_Dw_F</td>
<td>TTGGCAGTCGACTGAGAGACTG ATGGCCTT</td>
</tr>
<tr>
<td>RgpA_Dw_R</td>
<td>TACGCCAAGCTTGGAGAGCAGCAG ATTG</td>
</tr>
<tr>
<td>Tet_BamHI_F</td>
<td>TCAGGATCCACAACGAAATTATCCT CTTAAC</td>
</tr>
<tr>
<td>Tet_Sall_R</td>
<td>CGAGTCGACTGCAAGTTCTAATG CTTC</td>
</tr>
<tr>
<td>puc_EcoRI_R</td>
<td>ACCGAATTCCTGGCCGTCGT</td>
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Next, the RgpB deletional plasmid (pRgpBdel-erm) was obtained by the truncation of pRgpBall-erm plasmid using the PCR based Site-directed, ligase-independent method (SLIM) [145] with primers listed in Table 2. With this approach, only the small fragment of 3' (154 bp) of \( \text{rgpB} \) CDS was preserved on the pRgpBdel-erm plasmid. The \( \Delta \text{RgpB} \) strain was obtained in the homologous recombination event: the plasmid was elecroporated into the \( P. \text{gingivalis} \) W83 strain and the positive

Table 2: Primers used for construction of plasmids for \( P. \text{gingivalis} \) mutagenesis.
recombinant clones were selected with 5 µg/ml erythromycin. Proper recombination was verified by sequencing.

**Generation of *P. gingivalis* gingipains Q mutants**

Mutagenesis of each gingipain required a dedicated master plasmid. For RgpB studies the pRgpBall-erm construct was used. The Q1N mutation (replacement CAG codon into AAC) was incorporated using the SLIM method (resulting in the pRgpBallQ1N-erm plasmid), the Q25A mutation (replacement of the CAG codon into GCT) was introduced by the QuikChange method (Stratagene) (generating pRgpBallQ25A-erm), all sequences of applied primers are listed in Table X in the RgpBQ1N section. These plasmids were introduced into the *P. gingivalis* W83 RgpA-C strain lacking the whole *rgpA* gene [142] by the electroporation and the recombined clones were selected with 5 µg/ml of erythromycin. The obtained strains were partially sequenced and named ΔRgpA/RgpBQ25N and ΔRgpA/RgpBQ25A, respectively.

For RgpA mutagenesis, a master plasmid pNRgpA-tet was engineered. Two DNA fragments were amplified from *P. gingivalis* genomic DNA. The upstream 915 bp fragment consisting of sequence directly adjacent to the RgpA promotor was amplified with primers RgpA_Up_F and RgpA_Up_R). A downstream 2835 bp fragment, comprising the 5’ sequence of the RgpA gene together with 388 bp of its proposed promotor was amplified with primers RgpA_Dw_F and RgpA_Dw_R.
The tetracycline (tetQ) resistance cassette was amplified from the pT-COW plasmid [146] with primers Tet_BamHI_F and Tet_SalI_R. The backbone, the pUC19 plasmid, was linearized by PCR reaction with primers puc_EcoRI_R and puc_HindIII_F. All four amplified fragments were combined in the single step reaction by the method described by Gibson [144]. The Q24N mutation (replacement cag codon into aac) was incorporated to the construct with the SLIM method. Sequences of applied primers are listed in Suppl. Table 1 (RgpAQ24N section). This plasmid was introduced into the \textit{P. gingivalis} W83 \Delta RgpB strain lacking the whole rgpB gene by the electroporation and the recombined clones were selected with 1 µg/ml of tetracycline. Obtained strain was partially sequenced and named RgpBdel/RgpAQ24N. The non-mutated master plasmid was also introduced into the \Delta RgpB strain \textit{P. gingivalis} W83 and the unaffected expression and activity of RgpA was observed.

For Kgp mutagenesis, the pNKgp-cep master plasmid was created in a similar manner. First, two fragments adjacent to the start of the hypothetical Kgp promotor were amplified from the genomic DNA, the 809 bp upstream fragment with Kg_Up_F and Kg_Up_R primers, while the 3271 bp downstream fragment with Kg_Dw_F and Kg_Dw_R primers. The beta-lactamase gene \textit{cepA} was amplified with primers CepA_F and CepA_R from template synthetized by the Life Technologies based on the sequence deposited under AAA21538.1 number (Gene Bank). The pUC19 plasmid was linearized with primers pUC_Sphi_R and pUC_BamHI_F. The Q20N mutation (replacement caa codon into aat) was
incorporated to the construct with the SLIM method. Sequences of used primers are listed in Suppl. Table 1 (KgpQ20N section). This plasmid was introduced into the wild type *P. gingivalis* W83 by the electroporation and the recombined clones were selected with 2 µg/ml of ampicillin. Obtained strain was partially sequenced and named KgpQ20N. As a control, the non-mutated master plasmid was also electroporated into the *P. gingivalis* W83 strain and the unaffected expression and activity of Kgp was observed.
**Fig 3:** *P. gingivalis* mutant strains. A) schematic representing RgpBdel-erm. B) RgpA-tet C) Kgp-cep. Schematic representation of steps undertaken to generate *P. gingivalis* mutant strains.

**Gingipain activity assay**

The amidolytic activities of Rgp and Kgp enzymes were assessed by the hydrolysis of the chromogenic substrate benzoyl-L-arginine-p-nitroanilide (BApNA) and carboxybenzoyl-L-lysine p-nitroanilide (zKpNA; Novabiochem, Germany), respectively. In a 96-well format, 20-μl samples were preincubated in assay buffer
(200 mM Tris-HCl, 100 mM NaCl, 5 mM CaCl₂ [pH 7.6][71], supplemented with fresh L-cysteine to 10 mM) for 2 min prior to the addition of 1 mM substrate in a total volume of 200 μl. For activity measurement of Sarkosyl-treated membrane fractionations (see Materials and Methods), a 0.125 mM concentration of a synthetic arginine substrate pyro-glutamyl-glycyl-L-arginine-p-nitroanilide (pyroEGRpNA; Pharmacia-Harper, Uppsala, Sweden) was used instead of BApNA due to precipitation of the BApNA substrate in the presence of the Sarkosyl detergent. The presence of 0.1% Sarkosyl detergent in the assay did not affect the rate of substrate hydrolysis as determined with purified RgpB (data not shown). The rate of formation of p-nitroanilide was measured at 405 nm using a SpectraMax Plus spectrophotometer (Molecular Devices Inc., CA). For ease of comparison between mutants and statistical analyses of independent repetitions, activity units were defined as the total activity present in the RgpB⁺ control mutant culture equaling 100 U for culture partitioning studies, the total activity in the RgpB⁺ control mutant cells equaling 100 U for cellular fractionation studies, or the total activity in the RgpB⁺ control mutant membranes equaling 100 U for membrane fractionation studies.

**In vivo QC specific inhibition**

Each inhibitor was dissolved in DMSO at 10mM and 100mM concentrations. Five ml of a stock culture of *P. gingivalis* was equilibrated to OD₆₀₀ = 0.1 in 3 separate sealed tubes of 5ml of eTSB media (experiment repeated in triplicate). These tubes
are of the correct diameter to be read using the spectrophotometer cuvette reader. At inoculation either inhibitor concentration of 100µM or 1mM final concentration (DMSO content ≤ 1%) was added or equivalent DMSO ≤ 1% alone. Inoculated cultures of *P. gingivalis* were subsequently placed in an anaerobic chamber. Samples consisting of 1.0ml of bacteria were retrieved at 2, 5, and 8 hours post-inoculation. Samples were then centrifuged, supernatant was decanted, cell fraction was washed twice with PBS to remove unbound excess inhibitor. Bacterial pellet re-suspended in 300µl PBS and lysed by sonication (3 pulses, 5 s/pulse). Eighty µl of lysate used for continuous spectrophotometric assay of glutaminyl cyclase as previously described.

**Affinity purified anti-QC Antibodies**

CNBr-activated Sepharose 4B (GE Healthcare Lot # 10039621) was prepared according to manufacturer’s specifications. CNBr-activated Sepharose 4B is a pre-activated resin using the cyanogen bromide method, which couples antibodies or other large proteins containing -NH₂ groups to the Sepharose media, without an intermediate spacer arm. rPgQC was dissolved in coupling buffer (0.1 M NaHCO₃ containing NaCl 0.5 M) at 5mg/ml of CNBr gel. CNBr beads were washed and swelled on a sintered glass filter. rPgQC protein solution mixed with swelled gel suspension in an end-over-end mixer for 2 hours at room temperature or overnight at 4°C. Remaining active groups were blocked by addition of blocking agent 1.0 M ethanolamine or 0.2 M glycine at approximately pH 8.0 (2 hours at room
temperature or overnight at 4°C). Excess protein was washed away by at least 3 alternating washes of coupling buffer and acetate buffer (0.1 M pH 4.5 containing 0.5 M NaCl). Blocking agent was removed with 3-4 washes of coupling buffer followed by storage buffer (PBS containing preservative NaN₃). Serum passed through 0.45µm filter prior to application to the column. Serum was loaded into the column, column washed with PBS until A₂₈₀ reached baseline. The antibody bound to the column is eluted with 0.1M Glycine-HCl, pH 2.8. Fractions collected into tubes containing 1.0M Tris pH 8.0, to neutralize the eluted antibody. Antibody fractions were pooled and dialyze vs. PBS and finally concentrated. ELISA was run on serum, flow-thru, and eluate (Fig. 4).
**Fig 4. Specificity of anti-QC antibodies.** A 2-fold serial dilution afterwards was continued until no detection using both Western Blot and ELISA, Western blot (left panel) and Elisa (right panel). ELISA titration was preformed using clear bottom 96-well plate. Affinity purified QC concentration was determined by BCA assay at 1/10 dilution to be =8.4mg/ml. Concentration of QC was used @ 0.84mg/ml for first dilution (1/512). A 2-fold serial dilution afterwards was continued until absorbance was no longer detectable at 450nm wavelength.

**SDS-PAGE and Western Blots**

Samples were analyzed using established protocols ([133]). Samples were first boiled in non-reducing SDS-PAGE sample buffer containing 2mM TLCK for 5 min to inactivate all gingipains prior to the addition of 1% β-mercaptoethanol and boiled for a further 5 min for complete denaturation. Samples were centrifuged briefly at 13,000 × g, 1 min to remove particulates and the supernatant separated on SDS-PAGE and gels were stained with Coomassie Brilliant Blue. For Western blot analysis resolved proteins were subsequently electrotransferred onto 0.22-µm-
pore-size nitrocellulose membranes and blocked in 2% BSA/PBS solution overnight. RgpB was detected using a 1:2000 dilution of anti-RgpB mouse mAb in TTBS (20mM Tris, 500mM NaCl, pH 7.5 supplemented with 0.1% Tween 20) for 3 h. Membranes were washed four times with TTBS before being probed for 2 h with a 1:2000 dilution of an alkaline phosphatase-conjugated rabbit anti-mouse polyclonal secondary antibody (Dako Cytomation, Denmark). Development was carried out using the AP Conjugate Substrate kit as per manufacturer's instructions (Bio-Rad Lab., CA, USA). The in vivo PgQC and recombinant PgQC was detected using a 1:100,000 dilution (1μg ml⁻¹) of affinity purified specific (previously described) anti-QC rabbit polyclonal antibody in 5% (w/v) skim milk/TTBS solution. Membranes were washed four times with TTBS before being probed for 1 h with a 1:40,000 dilution of an HRP-conjugated rabbit anti-mouse polyclonal secondary antibody (Sigma-Aldrich). Development was carried out using the chemiluminescent protocol according to manufacturer's specification.
Bacteroidetes SPI substrates typically have a Qln (Q) downstream of the SPI cleavage site

The starting point for this study was the observation that many secreted *P. gingivalis* proteins with a predicted signal peptide also had a glutamine (Q) residue immediately downstream. The generality of this observation was confirmed on a genome-wide basis using the batch version of SignalP (for gram-negative bacteria). These species designations were pooled into “superspecies” groups, with particularly well-defined Q-values, and an anomalously large number of predicted proteins with predicted signal peptides or type I signal peptides. In none of the analyzed species groups, the “superspecies” exceeded 15% with exception of *Cyanobacteria*, where 31% species were incompletely designated, often without clear signs that several species were pooled (judging from the number of proteins). Including the “superspecies”, 337 *Bacteroidetes* species, 13 *Chlorobi* species, 59 Spirochaetes species, 13 *Chlamydia* species, and 82 *Cyanobacteria* species were analyzed. The number of proteins with predicted signal peptides was above 50 in all analyzed species, guaranteeing that incomplete proteomes did not have a major influence on Q-values. The medium number of proteins with predicted signal peptides was 399 for the *Bacteroidetes*, 125 for *Chlorobi*, 163 for *Spirochaetes*,
128 for *Chlamydiae*, and 195 for Cyanobacteria. Removal of lipoproteins from the set of proteins with predicted signal peptides was a relatively "small correction". The median values of the number of proteins with predicted type I signal peptides per species were 299 for the *Bacteroidetes*, 104 for *Chlorobi*, 146 for *Spirochaetes*, 112 for *Chlamydiae*, and 161 for *Cyanobacteria*. As lipoproteins have a C after the SPII cleavage site, their removal always increased predicted species Q-values. A "Q" immediately downstream of the signal peptide was predicted in about half of the cases. In the remaining cases, the residue after the signal peptide was frequently a cysteine, suggesting that these proteins were lipoproteins/SPII substrates. We used LipoP to identify and remove these proteins from the set. In the remaining set, which should only contain SPI substrates, the fraction of proteins with a Q after the SPI cleavage site exceeded 60%. Additional manual checks, including checks with earlier versions of the SignalP program, suggested that the true fraction of SPI substrates with a Q directly after the cleavage site may be even higher (Fig 5).
Fig. 5: Q-Value distribution in *Porphyromonas* - Fraction of predicted secreted proteins with a glutaminyl residue immediately downstream of the SPI cleavage site (Q-value) for different *Porphyromonas* species. Light gray bars indicated predictions on SignalP alone, while the dark bars indicate the predictions after removal of predicted lipoproteins (identified by LipoP).

The enrichment of glutamine downstream of the SPI cleavage site did not appear to be specific for the proteins of a particular cellular compartment. CELLO [147] predictions identified SPI substrate proteins in the inner membrane, the periplasm, the outer membrane, and the extracellular space. In all compartments, the fraction of SPI substrate proteins with a Q immediately downstream of the SPI site was
48% or higher (Table 3), clearly indicating that Q enrichment was not characteristic for proteins of a specific compartment. Placement of some SPI client proteins in the cytoplasm by the CELLO server suggests that some predictions are in error. Even with this reservation, Q enrichment does not appear to be characteristic for SPI client proteins in a particular compartment. This conclusion was further strengthened by the inspection of protein lists.

Table 3: CELLO Predictions - Proteins with or without Q residue in P1’ downstream of a type I signal peptide according to their localization, predicted by CELLO program.

Recombinant *P. gingivalis* PG2157 has QC activity and resides in the inner membrane

The high frequency of newly exposed Q residues in SPI substrates in *P. gingivalis* and the previously reported detection of 7 *P. gingivalis* proteins with N-terminal pyroglutamate [134] suggested that glutamine cyclization might not be
spontaneous, but enzymatically catalyzed. A BLASTP query of the *P. gingivalis* proteome with the human QC sequence suggested that PG2157 (also called PG_RS09565) may have QC activity. The recombinant protein (without a signal peptide) did not exhibit aminopeptidase activity on any of the commercially available substrates of general formula NH$_2$-L-Xaa-pNA or NH$_2$-L-Xaa-AMC. However, it efficiently converted the fluorogenic substrate L-glutaminyl-AMC into its respective pyroglutamic acid derivative ($K_m = 0.473 \text{ mM}$, $k_{cat} = 0.356 \text{ s}^{-1}$, $k_{cat}/K_m = 1.34 \text{ mM}^{-1}\text{s}^{-1}$). We therefore refer to PG2157 (PG_RS09565) as PgQC and to the recombinant version of the protein as rPgQC. SDS-PAGE of purification of rPgQC shown in Fig. 6.
Fig. 6: Purification of rPgQC. SDS-Page of different stages of expression and purification of recombinant *P. gingivalis* QC in *E. coli* BL21. Lane 2: Initial bacterial lysate applied to column. Lane 3: Flow through of unbound proteins. Lane 4: The fraction of bound recombinant protein plus the binding domain containing PSP cleavage sites and HIS-tagged region. Lane 5: Processing of the binding domain from the mature protein. Lane 6/7: Separation of the binding domain from the rPgQC. Lane 7: Final sample after reapplication on Ni-Sepharose column to further remove binding domain. Gel electrophoresis was completed using 10 µl of respective sample and run according to established protocol. (See Materials and Methods)
Table 4: rPgQC Enzyme Kinetics

<table>
<thead>
<tr>
<th>Species</th>
<th>H-Gln-AMC</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Km (mM)</td>
<td>$K_{cat}$ (s$^{-1}$)</td>
<td>$K_{cat}/Km$ (mM$^{-1}$s$^{-1}$)</td>
</tr>
<tr>
<td><em>Porphyromonas Gingivalis</em></td>
<td>0.473 ± 0.004$^a$</td>
<td>0.356 ± 0.02$^a$</td>
<td>1.34 ± 2$^a$</td>
</tr>
<tr>
<td>Human QC</td>
<td>51.3 ± 3</td>
<td>5.4 ± 0.1</td>
<td>98.0 ± 2$^b$</td>
</tr>
<tr>
<td>Human IsoQC</td>
<td>0.017 ± 0.001$^y$</td>
<td>1.07 ± 0.03$^y$</td>
<td>63 ± 6$^y$</td>
</tr>
<tr>
<td>Murine QC</td>
<td>0.048 ± 0.003$^δ$</td>
<td>6.0 ± 0.2$^δ$</td>
<td>125 ± 4$^δ$</td>
</tr>
<tr>
<td>Murine IsoQC</td>
<td>0.030 ± 0.006$^ε$</td>
<td>6.98 ± 0.35$^ε$</td>
<td>322 ± 27$^ε$</td>
</tr>
</tbody>
</table>

Sequence analysis of PgQC revealed a canonical signal peptide with the typical lipobox (Leu-Ser-Ala-Cys), suggesting that PgQC is a lipoprotein. Lipoproteins are translocated across the inner membrane via the Sec system, and initially anchored to the inner membrane by covalent attachment of a lipid anchor to the cysteine residue, followed by signal peptide cleavage, and typically second N-acetylation. Some, but not all lipoproteins are subsequently transferred from the inner to the outer membrane. Therefore, we expected that PgQC should be anchored in the inner or outer membrane.
In order to determine QC localization experimentally, cell extract (CE) of *P. gingivalis* in late exponential/early stationary phase of growth was fractionated into cytoplasm and periplasm (CP), total membranes (M), outer membrane (OM), and inner membrane (IM). The purity of the membrane fractions was verified by the exclusive presence of the biotin-containing 15-kDa biotin carboxyl carrier protein (AccA alias MmdC or PG1609) and gingipains in the IM and OM, respectively [136] (data not shown). QC activity of the fractions was then measured using the enzyme-coupled assay already used previously to demonstrate the activity of the recombinant enzyme. QC activity was found in the IM and in fractions containing IM (CE and M) but not CP and OM (Fig. 7A) clearly indicating that PgQC is anchored in the inner membrane. This localization was further confirmed by Western blot analysis of enriched subcellular fractions using rabbit polyclonal antibodies anti PgQC (Fig. 7B).
Fig. 7: PgQC is an Inner Membrane Protein. *P. gingivalis* in the early phase of growth was fractionated into sub-cellular fractions including, whole lysed cell extract (CE), cytoplasm (CP), membranes (M), outer membrane (OM), inner membrane (IM), and QC activity was measured in each sample results given in activity FU/s/µg. The QC activity in each fraction was determined with L-Gln-AMC as a substrate using a coupled assay with pyroglutamyl aminopeptidase as an auxiliary enzyme. The activity is shown as RF/s. The presence of PgQC antigen in fractions was determined by western blot.
Porphyromonas gingivalis QC is a zinc metalloenzyme

Previously, the metal ion content of human QC was unclear. Schilling et al. proposed that human QC is a metalloenzyme based on the inhibition by several metal ion chelators and the reactivation of apoenzyme by the subsequent addition of zinc ions [92]. However, Booth and co-workers reported lower than 0.3 molecule zinc ion per hQC molecule which was shown by mass spectrometry [120]. Because neither the accessory enzyme for the reaction, pyroglutamyl aminopeptidase, nor glutamic acid dehydrogenase, is inhibited by imidazole within the concentration range used, both the fluorimetric and spectrophotometric assay were well adapted for our purposes. The fluorimetric activity assay data (Table 5) revealed inhibition by imidazole. Imidazole completely blocks substrate conversion by binding in the active site. Inhibition of the enzyme occurs by removal and chelation of the metal ion required for catalytic activity, leaving an inactive apoenzyme. 1,10-Phenanthroline has been shown to mainly target zinc metallopeptidases and the inhibition of QC by 1,10-phenanthroline has been previously described [99]. EDTA has been shown under certain conditions to have an activating effect on QC catalysis and our data confirms that small amounts display an activating effect. It has previously been suggested that inhibition by phenanthroline is not due to metal chelation [94]. Also, in addition to being inhibited by 1,10-phenanthroline, P. gingivalis QC-catalyzed substrate cyclization was abolished in presence of dipicolinic acid, another inhibitor of metalloenzymes. Both chelators inhibited QC in a concentration and time-dependent manner, i.e. initial activity that was already
inhibited was found to be further reduced after prolonged incubation with the compounds (Fig. 8). However, EDTA did not show remarkable inhibition of *P. gingivalis* QC until the concentration was increased almost to 2.0 M. *P. gingivalis* QC was almost completely inactivated addition of 5 mM 1,10-phenanthroline. Schilling et al. have previously shown that repeated dialysis with chelator-free buffer, human QC activity was partially reactivated up to 50–60% but only in the presence of EDTA [94].

<table>
<thead>
<tr>
<th>Residual activity (%)</th>
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</thead>
<tbody>
<tr>
<td>0.1 mM</td>
</tr>
<tr>
<td><strong>Imidazole</strong></td>
</tr>
<tr>
<td><strong>Cysteamine</strong></td>
</tr>
<tr>
<td><strong>DTT</strong></td>
</tr>
<tr>
<td><strong>Glutathione</strong></td>
</tr>
<tr>
<td><strong>L-Cysteine</strong></td>
</tr>
</tbody>
</table>

Table 5. *Attenuation of rPgQC activity by reducing agents and imidazole.* Reactions were carried out at 30 °C in 0.06 M acetic acid, 0.06 M Mes, and 0.12 M Tris adjusted to the respective pH by the addition of NaOH or HCl. Data shown as residual activity set as 100% for given reaction without inhibitor. Data points are averaged triplicates.
Fig. 8: Concentration dependent inhibition by 1,10-phenanthroline and EDTA
Percent residual activity of QC in the presence of either compound was determined directly after the addition or pre-incubation of QC with each respective reagent for 15 min at 30 °C

To confirm effect of ionic strength on the activity of rPgQC, activity assay buffer was supplemented with differing concentrations of NaCl and KCl. rPgQC activity saw a sharp drop in activity with the addition of ion salts, but then activity tapered off to around 50% activity as the concentration of ion salts increased dramatically. (Fig. 9)
Fig. 9: Effect of differing concentrations of ionic salts on rPgQC activity. Activity using continuous fluorimetric assay displayed as percent activity of QC without ionic salts present in the activity assay buffer.

Sensitivity of the QC reaction to metal ions was also probed. Using the continuous fluorimetric assay, a reaction mixture was pre-incubated with the respective ion salt. Out of tested divalent cations only Cu$^{2+}$ exerted strong inhibitory activity (Table 6). Other metal ions had only slight effect only at 10 mM concentration in the reaction mixture. Interestingly but not surprisingly, increasing Zn$^{2+}$
concentrations within the QC assays (0.1 mM and higher) considerably reduce QC activity, and this has been observed in previous studies [94, 123].

Table 6: Residual activity of rPgQC in the presence of competing metal ions.

<table>
<thead>
<tr>
<th>Metal Ion</th>
<th>0.1 mM</th>
<th>1 mM</th>
<th>10 mM</th>
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<tr>
<td>MgCl2</td>
<td>88</td>
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<td>71</td>
</tr>
<tr>
<td>CaCl2</td>
<td>93</td>
<td>82</td>
<td>61</td>
</tr>
<tr>
<td>CuSO4</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>BaCl2</td>
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<td>61</td>
</tr>
<tr>
<td>MnCl2</td>
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<td>13</td>
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<tr>
<td>ZnCl2</td>
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</tr>
<tr>
<td>FeCl3</td>
<td>88</td>
<td>77</td>
<td>18</td>
</tr>
<tr>
<td>CuCl2</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 6: Residual activity of rPgQC in the presence of competing metal ions.

Given as percent activity of rPgQC standard. Performed on 96-well plate in triplicate, experiment repeated in triplicate. Values represent mean of the different results.
Pyroglutamate is present in signal peptidase I substrates

In order to experimentally demonstrate pyroglutamate at the amino-terminus of SPI substrates, previously determined mass spectrometry data was re-analyzed for *P. gingivalis* outer membrane vesicles [73, 136]. Outer membrane vesicles (OMVs) are continuously shed. Their vesicle lumen (VL), vesicle membrane (VM) and vesicle surface (VS) fractions contain proteins that are derived from the periplasm, the outer membrane, and extracellular proteins anchored to the outer membrane surface, respectively [148]. Pyroglutamate was inferred from the mass of the identified peptide, which is 17 Da less than what the unmodified peptide, and also by the fragmentation (MS/MS) pattern of the peptide. The MS/MS spectra of these peptides indicate that the -17 Da modification is present near the N-terminus of each peptide, most consistent with pyroglutamate formation. Altogether 27 proteins, all putative SPI substrates, with N-terminal pyroglutamate were identified. Interestingly, no semi-tryptic peptides with N-terminal glutamine were found in the entire *P. gingivalis* dataset, suggesting widespread pyroglutamate formation, despite the incomplete evidence for the complete set.

In order to further confirm this conclusion, we compared sequences around the cyclization site (and thus the SPI cleavage site) for proteins that were experimentally identified, had a glutamine after the predicted SPI cleavage site, and for which evidence for pyroglutamate formation was either available or not.
The region upstream of the SPI cleavage site, normally not expected to influence QC, was included in the comparison in case SPI and QC may act together and QC preferences may be influenced by SPI preferences. However, we did not detect clear differences in the sequence logos of the two groups of proteins on either side of the critical Q residue, further supporting the conclusion that QC acts broadly and is not limited in its activity by specificity for residues adjacent to the substrate glutamine residue (Table 7). Also noted is the fact that proteins with experimental evidence for pyroglutamate formation at the N-terminus were present in all compartments of outer membrane vesicles, this would subsequently stand against a role in partitioning proteins between VL, VM or VS of OMVs or the equivalent periplasmic space, outer membrane and outer membrane surface that these proteins stem from (Table 7).
Table 7: Q-value. Proteins from these compartments stem from the periplasm, the outer membrane, and the outer membrane surface, respectively. The row “Total” indicates the number of localized proteins, the row “Q” the number of proteins with a Q exposed by SPI signal peptide cleavage (according to a prior manual annotation, not focused on pyroglutamate formation), the row “SIGNALP = Y” the number of proteins with signal peptide, and the row “SIGNALP = Y&Q” the number of such proteins exposing a glutamine after signal peptide cleavage. The row “pGlu detected” identifies the number of proteins with experimentally verified pyroglutamate. In very few instances, a pyroglutamate was detected in a protein not predicted to expose a glutamine after signal peptide cleavage due an erroneous SIGNALP prediction.

<table>
<thead>
<tr>
<th></th>
<th>P. gingivalis</th>
<th></th>
<th>T. forsythia</th>
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<tr>
<td></td>
<td>VL</td>
<td>VM</td>
<td>VS</td>
</tr>
<tr>
<td>Total</td>
<td>27</td>
<td>79</td>
<td>30</td>
</tr>
<tr>
<td>Q</td>
<td>19</td>
<td>36</td>
<td>24</td>
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<tr>
<td>SIGNALP = Y</td>
<td>19</td>
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<td>20</td>
</tr>
<tr>
<td>SIGNALP = Y&amp;Q</td>
<td>10</td>
<td>19</td>
<td>16</td>
</tr>
<tr>
<td>pGlu detected</td>
<td>7</td>
<td>11</td>
<td>6</td>
</tr>
</tbody>
</table>
Q downstream of the SPI cleavage site affects RgpA, but not RgpB and Kgp secretion

In order to confirm that pyroglutamate formation does not affect secretion, we needed a model system in which pyroglutamate formation does not influence protein stability by controlling resilience against aminopeptidases. Gingipains RgpA, RgpB and Kgp, the major proteolytic virulence factors of P. gingivalis, are good model systems in this respect. The proteins are initially expressed as preproproteins, and their type I signal peptides are then cleaved upon import into the periplasm, exposing N-terminal glutamine residues. The CTD-domain containing proteins are then exported further by the type IX secretion system (T9SS). Upon secretion from the periplasm to the outer membrane surface, the pro-regions are rapidly degraded, leaving only the mature forms that no longer contain the expected pyroglutamate N-terminal residue. This is an asset for comparing protein activity independent of issues of protein stability, but it unfortunately also prevents detection of the pyroglutamylation of the pro-region in wild-type P. gingivalis strains. Nevertheless, we expect pyroglutamate formation based on its widespread occurrence (see above), and also because proRgpB is blocked for Edman degradation in a type IX secretion (T9SS) mutant that retains non-degraded proRgpB in the periplasm.

We used homologous recombination to construct P. gingivalis W83 strains expressing RgpAQ24N, RgpBQ25N, RgpBQ25A and KgpQ20N, in ΔRgpB,
ΔRgpA and wild-type backgrounds, respectively. Due to overlapping RgpA and RgpB specificities (both enzymes cleave after arginine residues), RgpA activities had to be compared in ΔRgpB background, and vice versa, whereas Kgp activities could be compared in a wild-type background. Mutation of the RgpA, RgpB or Kgp glutamine after the SPI cleavage site did not alter *P. gingivalis* growth or the extracellular activity of prolyl tripeptidyl peptidase secreted by *P. gingivalis* independent of T9SS. Gingipain activity was assayed in full cultures and the cell-free culture medium from cultures grown to the mid-exponential (OD$_{600\text{nm}}$ = 0.6-0.8), late exponential/early stationary (OD$_{600\text{nm}}$ = 1.2-1.4) and stationary (OD$_{600\text{nm}}$ >2.0) phase of growth. We assayed either total extracellular activity, or separately the activities associated with the outer membrane surface and with the culture medium. In all cases, activities increased strongly over time from mid-exponential to late stationary cultures. Mutation of the Q after the SPI cleavage site however, had surprisingly variable effects on the different gingipains.

The Q24N mutation in RgpA reduced overall activity several-fold, whereas the equivalent substitution Q25N (and also Q25A) in RgpB and Q20N in Kgp lacked significant effect, irrespective of whether cells were assayed in mid-exponential, late exponential/early stationary or late stationary phase (Fig. 10).

For all three gingipains, most of the activity was cell-associated, with only a minor contribution from protein in the medium. For RgpA, the Q24N decreased cell
Porphyromonas gingivalis strains were grown in at least six independent cultures to mid-exponential (OD600 in the range 0.6–0.8), late exponential/early stationary (OD600 in the range 1.4–1.6) and late stationary phase (OD600 > 2) of growth then adjusted to the same OD600 of 0.6, 1.4, and 2, respectively. Gingipain activity was measured in whole cultures with appropriate substrates. One-way ANOVA tests were carried out separately for the three gingipains (degrees of freedom RgpA: 5,30; RgpB: 8,45; Kgp: 5,48). Significance was judged applying the Bonferroni correction for multiple hypotheses (conservatively assuming all against all comparisons, even though only comparisons between wild-types and mutants are of interest) for a p = 0.05 threshold. The ** symbol indicates p < 0.001, the **** symbol p < 0.0001.

associated activity several fold, in all phases of growth. Activity in the medium was also significantly reduced in late stationary phase, the only growth stage with more than a marginal contribution from protein in the medium to the overall activity. In contrast, for RgpB, both the Q25N (and also Q25A) mutations altered the activity
associated with cells and in the medium at most insignificantly. There was also no redistribution of activity between cells and medium. For Kgp, the Q20N substitution had an unexpected effect. Although the overall activity was not significantly altered, Q substitution in Kgp increased activity in the medium at the expense of activity associated with cells, especially in mid-exponential phase, and to a lesser extent also in late exponential/early stationary phase.

We suspect that the diverse effects of Q substitution must be related to complicated posttranslational processing of Kgp and RgpA leading to an assembly of large multidomain complexes of the catalytic and hemagglutinin-adhesion domains derived from initial polyproteins anchored into the OM via A-LPS [149] rather than \textit{per se} translocation of the OM.

**QC is Essential for \textit{P. gingivalis} Survival**

Creation of non-polar QC knock-out strains in \textit{P. gingivalis} was an untenable goal due to apparent lethality of absence of QC activity. \textit{P. gingivalis} QC (PG_2157) lies within an operon containing genes PG_2157, PG_2156, and a HemG (PG_2158), and in initial attempts downstream effects were present in the gene replacement (Fig. 11). Subsequently we attempted construction of non-polar knock out strains, without disrupting the operon but it was evident that gene PG1885 is essential for survival of \textit{P. gingivalis} bacteria. Partial success was
achieved with only one survivable mutant in which erythromycin was successfully inserted between PG_2157 and PG_2158.

Fig. 11: Structural Schematic of the *P. gingivalis* operon containing QC. Gene coding sequence PG_2157 (QYT). PgQC lies just prior to PG_2156 and PG_2158 and non-polar disruption of PG_2157 was lethal.

Following these revelations, we shifted focus into examining and manipulating the possible QC targets that were identified during the SignalP QC distribution data.

*P. gingivalis* growth attenuation by PgQC Inhibition

After assessing the effects of non-specific inhibitors of QC activity both *in vitro* and *in vivo*, specific inhibitors were produced to experimentally determine the effects of reduced PgQC activity on bacterial growth. With the assistance and cooperation of Probiodrug in Germany, banks of specific inhibitors were screened against purified rPgQC. Promising candidates were then analyzed for inhibition on QC
activity. The initial inhibitor was LL29. Inhibitor LL29 was dissolved in DMSO into 100mM concentration and incubated with fresh *P. gingivalis* liquid culture. 1.5ml of liquid culture of *P. gingivalis* were lysed, and remaining LL29 inhibitor was washed with PBS and QC activity was recorded using fluorimetric assay. In these cell lysates cultures that were administered 100mM LL29 showed a time-dependent competitive inhibition of PgQC. In addition to the in vitro effects of LL29 on PgQC, corresponding 1.5 mL cultures of early-log phase *P. gingivalis* were incubated overnight to examine the inhibitors’ effect on *P. gingivalis* growth. After 24-hour incubation in anaerobic chamber, the OD$_{600}$ of cultures was analyzed. LL29 inhibitor incubated cultures showed a decrease in absorbance equivalent to 33% cell density decrease. Initially we suspected that the cell permeability may be a complicating factor or extracellular inhibitor could be degraded by cell factors. To effectively rule out inhibitor penetration into bacteria cells, these experiments were repeated in triplicates but concentrations of LL29 were increased 10 fold to 1mM and compared to initial 100mM concentration. (Fig. 12)
Fig. 12: LL29 Inhibits P. gingivalis growth. 5 ml of a stock colony of P. gingivalis equilibrated to OD$_{600}$ = 0.1 in 3 separate sealed tubes of 5 ml of eTSB media in triplicate. At inoculation 1mM inhibitor 084 final concentration (DMSO ≤ 1%) was added or equivalent DMSO ≤ 1% alone was added. Inoculated colonies of P. gingivalis were subsequently placed in an anaerobic chamber. Statistical difference (p<0.0001) shown between DMSO and 1mM LL29 are demonstrated with ****.

What is striking is the group with 1mM LL29 showed a remarkable decrease in QC activity along with a 50% decrease in total liquid culture density (Fig 12). 24-hour growth curves were plotted to demonstrate LL29’s effect over time (Fig. 12B). Absorbance reading were taken at 1-hour intervals over a 28-hour period and plotted to display accurate growth curves. These showed that the cultures with 1mM LL29 only reached an absorbance of less than half of the control group (DMSO alone). RgpA-null strain cultures were pre-incubated with LL29, pelleted, washed three times with PBS and then resuspended in fresh eTSB media to
removed trace amounts of excess LL29 not taken up by the bacteria. This eliminated excess inhibitor from the reaction. 100mM LL29 demonstrated time-dependent inhibition of W83 in vivo QC activity. This inhibition was more pronounced at 1mM concentration (Fig. 13).

Fig. 13: PgQC in vivo inhibition by LL29. Analysis of QC activity in vivo W83 P. gingivalis. LL29 had a time-dependent inhibitory effect on QC activity at final concentration 100mM. Whole cell activity was measured after cell lysis. Statistical difference (p<0.0001) shown between DMSO and 1mM LL29 are demonstrated with ****.

Samples were then taken at 2, 5, and 8-hour marks and whole cell gingipain activity assays were completed. Using the data stated above, further screening of potential specific inhibitor produced 5 additional possible candidates. Inhibitors: 019, 071, 073, 101, and 084 were roughly screened using in vitro PgQC assays. Each inhibitor showed significant attenuation of PgQC activity but only 084 completely obliterated QC activity (Fig. 14). Previously, each respective inhibitor was incubated within the reaction mixture prior to initiation of the assay. In the case of
084, we wanted to determine if the inhibitor had a bacteriocidal or bacteriostatic effect, so the growth curve was repeated with addition of 1mM 084 at mid-logarithmic bacterial growth (Fig. 14F). After addition of 084, bacterial cell density decreased slowly up to 24 hrs after which it remained static. This suggests some mixed bacteriostatic/bacteriocidal killing of *P. gingivalis* after addition of inhibitor 084 at 1mM final concentration.
**Fig. 14: 24-hour growth Inhibition of *P. gingivalis* W83.** 5 ml of a stock colony of *P. gingivalis* equilibrated to OD<sub>600</sub> = 0.1 in 3 separate sealed tubes of 5 ml of eTSB media in triplicate. At inoculation 1mM inhibitor 084 final concentration (DMSO ≤ 1%) was added or equivalent DMSO ≤ 1% alone was added. Inoculated colonies of *P. gingivalis* were subsequently placed in an anaerobic chamber. Absorbance was measure at each hour for 24 hours. Inhibitor 084 had the most pronounced effect of inhibition *P. gingivalis* growth.
High Q-values are typical for Bacteroidetes species, but not related phyla

In the following, we call the fraction of signal peptidase I substrates that have a glutamine immediately downstream of the SPI cleavage site the Q-value. The high Q-value is not unique for P. gingivalis but is shared with other bacterial species. LipoP corrected SignalP predictions for various Porphyromonas species suggest Q-values between 59% (for P. somerae) and 77% (for P. macacae). Manual inspection of automatic prediction suggested that even the high Q enrichment values are still underestimated due to the mis-prediction of some signal peptide cleavage sites.

Porphyromonas species belong to Bacteroidetes, which in turn are placed in the FCB superphylum containing the Fibrobacteres, Chlorobi, and Bacteroidetes. In the Bacteroidetes group, 332 of 334 species (i.e. > 99%) had predicted Q-values above 48%. Averaged across species, the Q-value was 71%, with a standard deviation for the variation between species of around 10% percent (outliers included). The two low Q-value outlier species (Bacteroides pectinophilus, 4% and Candidatus cloacimonas 24%) had a suspiciously low number of predicted proteins with signal peptides (21 and 83, compared to typically 300 in this group), suggesting a possible problem with sequencing or annotation rather than a genuine difference. CELLO predictions for T. forsythia confirm the conclusion for P. gingivalis that the enrichment of glutamine residues directly after the SPI
cleavage site is not specific for proteins of a particular cellular compartment (Table 3). Across the entire FCB superphylum, Q-values were not consistently high. For most sequenced Chlorobi species, the predicted Q-value was below 9%. Three outliers were found in this group in little characterized bacteria (annotated as Chloroherpeton thalassium, Q-value 55%, Chlorobi bacterium, Q-value 63% and Chlorobium sp., Q-value 67%). Outside the Bacteroidetes and Chlorobi, a continuum of Q-values was found, ranging from 2% (Chlorobaculum parvum) to 77% (Ignavibacterium album), with no obvious pattern.
Fig. 15: Q-values averages and Q-value distributions for different groups of bacteria. The Q-value was determined separately for every species in the groups. (A) Average Q-values for the species groups (species are given equal weight in the average), and the standard deviation of Q-values. Bacteroidetes have significantly higher Q-values than Chlorobi, Spirochaetes, Chlamydiae, or cyanobacteria (p < 1E-6 for comparisons of Bacteroidetes against any of the other species groups according to the Wilcoxon rank sum test). (B) Cumulative Q-value distributions. For identification of species groups, refer to the symbols in the right panel (near Q-value 1). The Q-values for Bacteroidetes species are higher than for the other species groups also judging from cumulative distributions (p < 1E-6 according to the Kolmogorov–Smirnov test). Species were classified as having an animal- or plant-type QC according to BLASTP searches using the human and A. thaliana QC.

We also determined the cumulative Q-value distribution for selected bacterial phyla (Fig. 15), thought to be relatively closely related to Bacteroidetes [150]. Predicted Q-values for Chlamydiae and Spirochaetes species were typically below 20% and thus much lower than in Bacteroidetes. Cyanobacterial species typically also had
lower Q-values. However, in a small fraction of cyanobacterial species (~5%), Q-values were not much lower than those of \textit{Bacteroidetes}. High Q-values were for example found for \textit{Scytonema millei} (40%), \textit{Leptolyngbya valderiana} (45%), \textit{Hassallia byssoidea} (61%), and \textit{Aphanocapsa montana} (67%) (Fig 15).

\textbf{\textit{Bacteroidetes} have orthologues of animal and plant QCs}

We searched complete bacterial proteomes in UNIPROT for orthologues of animal- and plant-type QCs. In a first step, we used the prototypical human and \textit{A. thaliana} QC sequences as queries. In a set of 1878 proteomes, we found 602 animal-type QCs, among them 402 (67%) in \textit{Bacteroidetes}. We also identified 991 plant-type QCs, among them 401 (40%) in \textit{Bacteroidetes}. As \textit{Bacteroidetes} account for less than 10% of the proteome data (and even less in the redundant set), it is clear that QCs, particularly of the animal-type, are enriched in the \textit{Bacteroidetes}. Orthologues of animal- and plant-type QCs tended to segregate according to phylogeny. Animal-type QCs were typically found in \textit{Bacteroidia}, plant-type QCs in \textit{Flavobacteriia}. As the number of candidate QCs was smaller than the number of proteomes, we attempted to enlarge the set of candidate QCs by carrying out BLASTP queries with representative sequences from \textit{Bacteroidia} and \textit{Flavobacteriia}, the two largest groups within the \textit{Bacteroidetes}. Starting from these sequences, and correcting for species duplicates, we identified 574 \textit{Bacteroidetes} species containing animal-type and 507 \textit{Bacteroidetes} species containing plant-type QC enzymes, but only 63 species containing both types of
enzymes (E-value threshold 1E-4), compared to 1540 species in the duplication corrected proteome dataset.

We also attempted iterated searches, using representatives of CD-HIT identified sequence clusters [151] to initiate additional searches. With this procedure, still more putative QCs were identified, but the concentration of hits in the Bacteroidetes phylum was reduced, likely because the set of putative QCs became contaminated by peptidases. Despite this complication, we can conclude from the simpler BLASTP searches that most and perhaps even all Bacteroidetes species have enzymes that could be suitable for glutamine cyclization. In the following, we focus on the candidate QC enzymes that can be demonstrated in a single BLASTP step with tight E-value threshold to be orthologous to the prototypical animal- or plant QC enzymes.

The orthologues of animal and plant-like QCs in Bacteroidetes have intact active sites

In order to assess the chances that the animal- and plant-type QCs in bacteria were active, we checked alignments for the presence of key active site residues. The prototypical animal QC is the human enzyme, hsQC. Its active site is built from E201 (involved in proton shuttling), D159, E202 (involved in binding the active site Zn^{2+} ion), and D248 (involved in both). In the Bacteroidetes orthologues of human QC, D159, E201 and D248 are highly conserved (>98%). In contrast, E202 is
conserved only in 9% of *Bacteroidetes* orthologues, and replaced by an aspartate residue almost all remaining cases (exceptions <1%). As aspartate and glutamate can both serve as Zn$^{2+}$ ligands, we suspect that this substitution may not compromise activity, or may even be required to accommodate slight changes in the overall protein structure compared to HsQC. This conclusion is supported by the observation that an aspartate is also present in the 202 position in the PgQC enzyme, which we have already shown to be an active QC enzyme.

The prototypical plant QC is the enzyme from *C. papaya*, CpQC. Its active site is not as well understood as the active site of hsQC, but it is thought that E69, N155 (probably involved in proton shuttling) and N155, K225 and Q24 (probably involved in stabilizing the oxyanion intermediate) play a role in catalysis. Among the *Bacteroidetes* orthologues, E69, N155 and K225 are strictly conserved. In contrast, Q24 was conserved only in 81% of cases, and replaced by a glutamate in the remaining cases. The same substitution occurs in many plant enzymes, and also in the experimentally studied bacterial *Xanthomonas campestris* QC (XcQC). In this special case, the (natural) glutamate variant is still active, albeit an order of magnitude less so than the engineered glutamine variant [91], suggesting that both glutamine and glutamate in the active site are compatible with activity, although not necessarily at the same level.

We conclude from this analysis that *Bacteroidetes* orthologues of animal and plant QCs are likely to be active enzymes. This is directly suggestive of QC activity plant
QC orthologues, and compatible with either QC or aminopeptidase activity for animal QC orthologues. Classification of the enzymes as lipoproteins (like PgQC) could strengthen the case for QC activity.

**Most Bacteroidetes orthologues of animal and plant QCs are predicted lipoproteins**

Among 401 orthologues of human QC in *Bacteroidetes*, 323 (~80%) were predicted lipoproteins, 59 (~15%) were predicted SPI substrates (~15%), and the remaining 19 (~5%) were predicted to be cytoplasmic. As orthologues of human QC are highly enriched in *Bacteroidetes*, relatively few were found in species not belonging to the *Bacteroidetes*. Among these, the fraction of predicted lipoproteins was much smaller. Only 38 (~34%) enzymes were predicted to be lipoproteins, 40 (~36%) were classified as SPI substrates, and remaining 34 (~30%) as cytosolic proteins.

Among 430 orthologues of *A. thaliana* QC in *Bacteroidetes*, 376 were (~87%) computationally classified as lipoproteins, 20 (~5%) as SPI substrates, and 34 (~8%) as cytoplasmic proteins. The predominance of predicted lipoproteins among plant-type QCs was much less pronounced when bacterial homologues of plant QC in general were considered. Among the 1000 bacterial proteins most similar to *A. thaliana* QC, predictions classified 475 (~48%) as lipoproteins, 338 (34%) as SPI substrates, and 175 (18%) as cytosolic proteins.
Sensitivity (true positive rate, recall) of the LipoP algorithm for gram-negative bacteria has been reported to be around 90% [140]. The fraction of bacterial QCs predicted to be lipoproteins in non-

*Bacteroidetes* species is much smaller, suggesting that not all are lipoproteins. In *Bacteroidetes*, the fraction of QC proteins predicted as lipoproteins comes close to the predicted sensitivity of the prediction algorithm. Thus, it appears likely that most if not all QCs in *Bacteroidetes* are lipoproteins, like the prototypical PgQC from *P. gingivalis*.

**Proteomic datasets confirm glutamine cyclization in several *Bacteroidetes* species**

In order to confirm widespread pyroglutamylation formation in *Bacteroidetes*, and not only *P. gingivalis*, we analyzed additional data from previously reported proteomic studies [73, 134-137]. In addition to the already discussed 27 proteins from *P. gingivalis*, the collated data identify 27 proteins in *Tannerella forsythia*, 13 in *Parabacteroides distasonis*, 8 in *Prevotella intermedia* and 7 in *Cytophaga hutchinsonii* with N-terminal pyroglutamate residue. N-terminal residues other than pyroglutamate were rare, as predicted from the bioinformatic studies.

Pyroglutamate was not detected at the amino-terminus of all SPI substrates that are predicted to expose an N-terminal glutamine residue after SPI cleavage, most likely due to incomplete coverage and not due to selective pyroglutamate
formation. As already reported for the *P. gingivalis* proteins, semi-tryptic peptides were never found to start with glutamine, even though internal tryptic peptides could be identified with N-terminal glutamine in both modified and unmodified states. Other circumstantial evidence also supports widespread rather than selective glutamine cyclization. We focused in particular on the *T. forsythia* data, which contained pyroglutamate evidence for the largest number of proteins in one species. Experimentally detected proteins (by any peptide, not necessarily a semitryptic peptide) with glutamine after the SPI site were partitioned into proteins with and without evidence for glutamine cyclization (Figure 16). As already reported for *P. gingivalis*, amino acids around the Q were similar in the two groups, supporting broad QC specificity.

**Fig. 16: Sequence downstream of Q in *T. forsythia* SPI proteins.** Comparison of sequence logos around Q downstream of the signal peptide for *T. forsythia* SPI proteins. Top – experimentally demonstrated or Bottom – undetermined Q cyclization in OMVs. The signal peptidase motif A-X-A upstream of the SPI cleavage sites is marked by the arrow. Only proteins with a Q after the predicted SPI site have been included in the comparison panel in the bottom.
In addition, QC activity was detectable in *T. forsythia* using the previous method for continuous fluorimetric QC activity in *P. gingivalis* (Fig 17).

**Fig. 17: QC activity in *T. forsythia*.** *T. forsythia* in the early phase of growth was measured using whole cell extract, and QC activity was measured in each sample results given in activity FU/s/µg. QC activity was determined in each fraction standardized to the volume of washed cells were suspended for sonication. The QC activity was determined with L-Gln-AMC as a substrate using a coupled assay with pyroglutamyl aminopeptidase as an auxiliary enzyme. *P. gingivalis* W83 strain used here for comparison.

Following this experimentally measured QC activity of a *Bacteroides* sp. Other than *P. gingivalis*, we decided to test the specificity of LL29 against the *T. forsythia* QC. LL29 demonstrated no inhibition of QC activity *in vivo* at any time point in *T.*
forsythia (Fig 18). This could be easily explained by substrate preference or variability in the active site structural motifs.

**Fig. 18: LL29 Does not inhibit TfQC.** *T. forsythia* in the early phase of growth was measured using whole cell extract, and QC activity was measured in each sample results given in activity FU/s/µg. QC activity was determined in each fraction standardized to the volume of washed cells were suspended for sonication. The QC activity was determined with L-Gln-AMC as a substrate using a coupled assay with pyroglutamyl aminopeptidase as an auxiliary enzyme. *P. gingivalis* W83 strain used here for comparison. Statistical difference according to one-way ANOVA (p<0.0001) shown between DMSO and 1mM LL29 are demonstrated with ****.

Pyroglutamate detection is not consistent for paralogue families in a single species. Pyroglutamate detection for a protein of a given bacterial species is also
not predictive for the orthologue in another species. In a first step, we used BLASTP (E-value threshold $10^{-9}$) to identify orthologous proteins in *T. forsythia* and *P. gingivalis* with experimentally verified localization. Only unique pairs were used, and proteins with more than one paralogue in either species were excluded. Among the 41 pairs, 9 and 3 were found to have a pyroglutamate at the N-terminal end. Based on a random association alone, one would then expect experimental pyroglutamate demonstration for both proteins of a pair in $(9 \times 3)/41 \sim 0.7$ cases. In fact, one such case was observed.

Together, the above observations are consistent with general glutamine cyclization, partially masked by incomplete mass spectrometry coverage (for example, due to low expression of some proteins, or because semi-tryptic peptides are too short or too long for efficient mass spectrometry detection). As already seen for *P. gingivalis*, pyroglutamate formation was not characteristic for proteins of a particular compartment, as *T. forsythia* proteins with pyroglutamate were also found in the vesicle lumen, in vesicle membranes, and on the vesicle surface of OMVs, which represent periplasmic, integral OM and cell-surface associated proteins, respectively.

**A model for pyroglutamate formation in proteins destined to the periplasm, the outer membrane, the outer membrane surface, or the medium**

Together, this body of data suggest a model for *Bacteroidetes* SPI substrates that reside in or transit through the periplasm. Their signal peptides are cleaved by SPI,
a lipoprotein with active site on the periplasmic face of the inner membrane. Because of the enrichment of glutamine immediately downstream of the SPI cleavage site, this reaction typically exposes an amino-terminal glutamine residue. QC, another lipoprotein, also with active site on the periplasmic face of the inner membrane, is then ideally positioned to catalyze the cyclization of the glutamine residue to a pyroglutamate residue. The cooperation between SPI and QC is apparently efficient, suggesting either direct interaction or joint anchoring in lipid domains, which we have not yet tested. Pyroglutamate formation occurs for proteins that remain in the periplasm, as well as for proteins that are transported further into or through the outer membrane. We suggest the term “Q-rule” to describe the finding that glutamines are enriched after SPI cleavage sites, and that these glutamine residues are cyclized to form N-terminal pyroglutamate residues in proteins that are destined to the periplasm or beyond (Fig. 19).
Fig. 19: Hypothesized Pathway for Pyroglutamate formation by QC.
Schematic representation of the pathway for pyroglutamate formation at the amino
terminus of proteins that transit to or through the periplasm. It is currently unclear
whether or not SPI and QC are physically associated. Pyroglumate formation has
not yet been tested for proteins of the inner membrane.

Importance of the Q-rule pathway

The Q-rule pathway seems to be biologically important, not only judging by the
number of proteins that are subject to the rule. *P. gingivalis* glutaminyl cyclase is
present in both virulent and avirulent strains [152]. According to an unbiased large
scale transposon mutagenesis screen [153], the enzyme is essential even in
laboratory culture conditions when *P. gingivalis* is not pitted against a host immune
system. It is currently unclear why the enzyme is essential. Our data speak against
a role of the modification in protein sorting. Given the host-associated lifestyle of
many *Bacteroidetes* species, it is possible that glutaminyl cyclization protects secreted proteins against host aminopeptidases (excluding of course the host pyroglutamate aminopeptidases). However, this model does not explain why the Q-rule applies to SPI substrates that remain in the periplasm or why the glutaminyl cyclase is essential for *P. gingivalis* in culture conditions, unless *P. gingivalis* has become unable to do without the Q-rule and now needs pyroglutamate formation for protection against its own periplasmic proteases as well. Further lending credence to this theory is the lethality of QC knock-outs.

**Future topics**

One of the lingering questions remaining is how the QC and SPI interact within the periplasmic space. It was not evaluated whether these are or are not in intimate contact or physically linked to one another, although their function is clearly linked. Another point of interest is perhaps although only downstream effects of gingipain secretion were evaluated in this project, *P. gingivalis* is fully capable of surviving without gingipain activity. Perhaps another essential protein or proteins processed through this system is blocked, thereby leading to cell death. Either way this system poses as a possible target for future studies and continued research.
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