




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The development of matrix-metalloproteinase responsive sensors for the machine-independent detection of oral inflammation†

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Periodontitis is a chronic inflammatory disease that affects an estimated 20–50% of the world's population and can lead to high follow-up costs for the healthcare system. The disease is closely associated with a dysbiotic, pro-inflammatory change of the oral microbiota. As the disease progresses, the inflammation affects the periodontium with subsequent tooth mobility and tooth loss as clinical outcomes. Early diagnosis and early treatment are vital and can leverage overall clinical success, resulting in possible preservation. Therefore, periodontal inflammation should be detected as soon as possible. Unfortunately, early stages of periodontitis are mostly asymptomatic and unnoticed by affected patients or only noticed at advanced stages, making diagnosis impossible without medical consultation. Ideally, highly intuitive diagnostic tools would be available to anyone, anywhere and anytime for personal check-ups. Therefore, we developed peptide sensors, that are cleaved by upregulated matrix metalloproteinases (MMP) as present in oral inflammation. This leads to the release of a denatonium-based flavor substance, providing a rapid diagnostic indicator for further professional clarification. Two distinct techniques were employed to enhance sensor specificity. A peptide sensor was developed for disease-specific protease (MMP) digest using previously published information and proteome wide screening techniques, followed by optimization against nonspecific proteases present in the oral cavity. Finally, the selected sensor was tested in patient saliva, and the sensor cleavage correlated with MMP concentration and activity. Furthermore, the sensors significantly differentiated patient pools and pools of healthy volunteers, and cleavage was linked to pocket probing depths, a clinical sign of periodontal diseases.

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Introduction

Chronic inflammations of the periodontium, such as gingivitis and periodontitis, are among the most common oral diseases worldwide. If left untreated, they can lead to tooth loss. This is accompanied by significant functional limitations and considerable impairment of aesthetics and

phonetics, leading to exclusion and other adverse effects on social life.^{1,2} Worldwide, periodontitis affects an estimated 20–50% of the global population and around 40% of the adult population in the US.^{3,4} The consequential costs of late or undetected cases for the healthcare system are high and will likely rise with an aging population.^{5,6}

Periodontitis is a chronic inflammatory process fueled by microbial dysbiosis, dental plaque formation, genetic predisposition, systemic diseases, smoking, or lifestyle.⁷ Progressing inflammation drives irreversible damage to the tooth-supporting tissues and, ultimately, tooth loss.⁸ Therefore, therapy aims to start in the early stages of the disease.⁹ The sooner inflammation in the oral cavity is detected and treated, the greater the chance of preventing the occurrence of irreversible damage and maintaining oral health into old age.

Self-diagnosis is impossible. The clinical dental examination includes measuring pocket probing depth, recession, bleeding on probing (BoP), tooth mobility, furcation, and radiographic findings of bone loss.⁹

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Unfortunately, periodontitis often starts symptom-free, stays unnoticed, and is only noticed in an advanced state by affected patients. Thus patient compliance, even where patients are entitled to use the offer of an annual dental check-up of their oral health, is often insufficient, and an early diagnosis can usually not be guaranteed.^{9,10} That gap defines the central challenge for this manuscript.

Saliva offers easy, non-invasive diagnostic access to multiple diseases.¹¹ A variety of methods have been described targeting saliva, *e.g.*, proteomics, point of care testing (POCT), or electronic tongues.^{12–14} While proteomics and electronic tongues provide broader information, ranging from information of whole salivary proteomes to phonetic interpretation of complex liquids, POCT is mainly focused on

defined analytes and simple readouts.^{12,15} Here, promising saliva biosensing platforms can quantify analytes like virus particles to screen for infectious diseases or address non-communicable diseases such as cancer by microRNAs.^{16,17} A well-known example of a US Food and Drug Administration (FDA)- and EU-approved PoC test kit that uses mouth rinse samples rather than saliva is the commercially available aMMP-8-based lateral flow immunoassay (*e.g.*, PerioSafe®, Implant Safe®, Oral Risk Indicator®).¹⁸ Using digital reader systems, quantifiability of the results and high sensitivity is possible, allowing the mapping of the different stages of periodontitis and the selection of appropriate therapeutic approaches.^{19,20} In previous studies, we could achieve good results in the release of the bitter flavor with the electronic

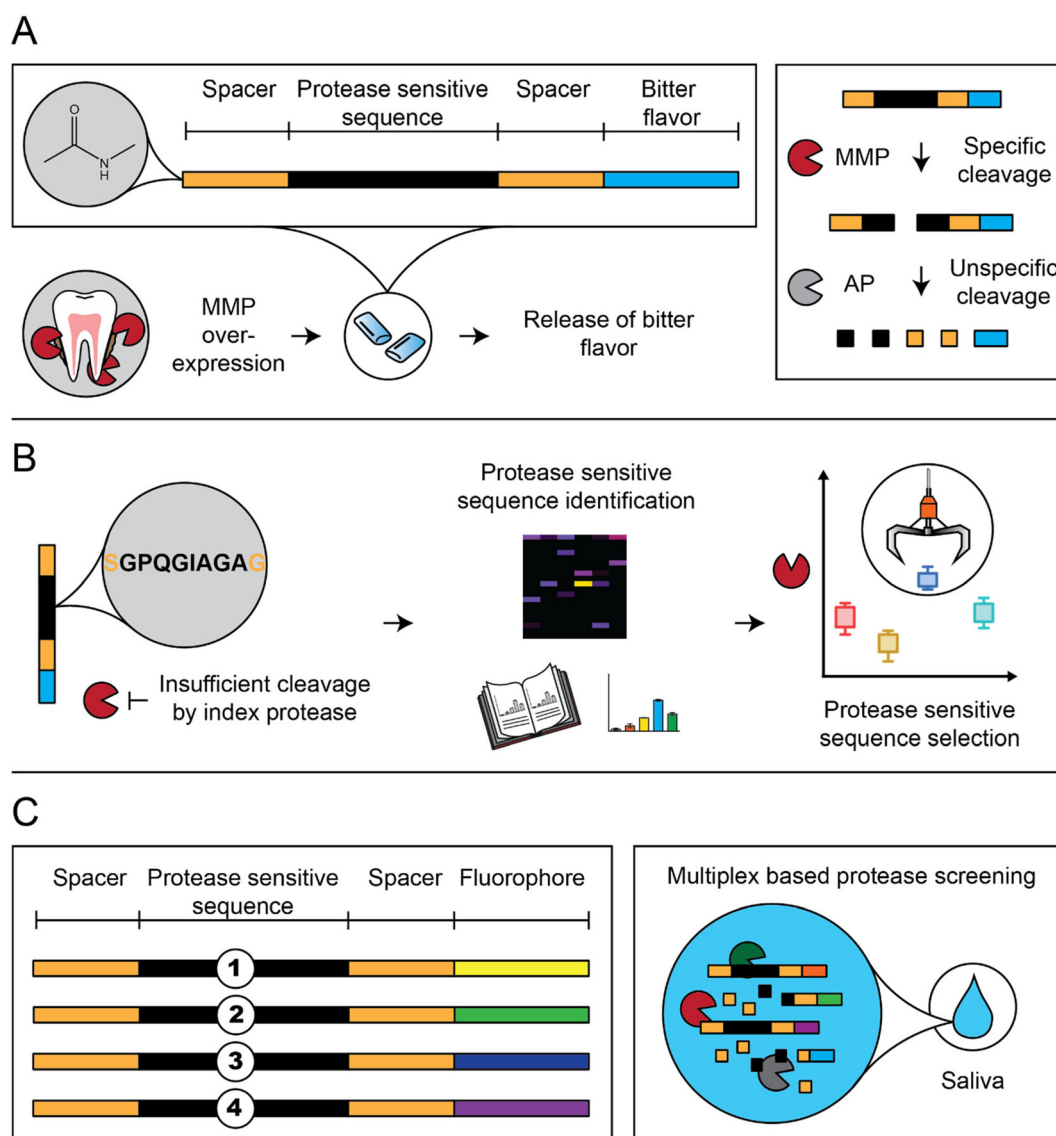


Fig. 1 (A) The design and principle of the peptide-based sensors. The index protease (MMP) specifically cleaves the protease-sensitive sequence (PSS), resulting in fragments. These fragments are further digested by unspecific aminopeptidase (AP), releasing a bitter substance. Acetylation of the sensor protects unspecific AP cleavage. (B) Cartoon of the steps for PSS identification and iterative improvement. (C) Final testing of selected sensors in saliva using multiplexing.



tongue using the precursor model, a particle-based sensor.²¹ With our new system presented here, we aim to achieve a simple, fast, and ubiquitous readout in complex fluids, *i.e.*, translating differences (healthy and periodontitis patients) in matrix metalloproteinase activity in saliva, a surrogate marker for inflammation, into a bitter substance and finally into taste.^{22,23} The taste is supposed to motivate the patients for professional consultation. Therefore, this system complements current pull strategies (patients comply with regular visits for check-ups) with push strategies (the sensory chewing gum pushes patients into the office).

The peptide-based sensors disintegrate by upregulated MMPs, surrogate markers of oral inflammation, and lead to the release of a bitter-tasting denatonium derivative (Fig. 1A).^{21,24} First, we screened our sensors with MMP-8, a collagenolytic protease correlating with oral inflammation, including periodontitis.^{25,26} We deployed proteome-wide search strategies (proteomic identification of protease cleavage sites; PICS) to screen for new protease-sensitive sequences (PSS; Fig. 1B).²⁷ In addition, selected sensors were *N*-acetylated to prevent unspecific proteolytic digestion by salivary aminopeptidases (AP). Finally, sensor selection was performed in patient saliva to include the full periodontitis orchestra (Fig. 1C).^{28–33}

Material and methods

Proteomic identification of cleavage sites (PICS)

Identification of protease cleavage sites was performed with minor changes according to PICS protocols described by Schilling *et al.*³⁴ Three different PICS libraries were generated, namely human embryonic kidney (HEK) 293 (ATCC-Number CRL-1573, ATCC, Manassas, VA,) cell based, *Escherichia coli* (*E. coli*) (Thermo Fisher Scientific, Waltham, USA) ¹⁴N and *E. coli* ¹⁵N. A detailed method section is provided in the ESI.†

Peptide synthesis

All peptides described in this manuscript were synthesized using fluorenylmethoxycarbonyl protecting group (Fmoc) based solid phase peptide synthesis, following established protocols using a polypropylene-reactor (MultiSynTech, Witten, Germany). A detailed method section is provided in the ESI.†

Liquid chromatography-mass spectrometry (LC-MS)

Peptide mass spectra were acquired *via* a Shimadzu LC-MS system (Shimadzu Scientific Instruments, Columbia, MD, USA) equipped with a DGU-20A3R degassing unit, a LC20AB liquid chromatograph using a Synergi 4 μ m fusion-RP column (150 \times 4.6 mm) (Phenomenex Inc., Torrance, CA), SPD-20A UV/vis detector, coupled with a single quadrupole LCMS-2020. Mobile phase A was 0.1% (v/v) FA in Millipore water. Mobile phase B was methanol with 0.1% (v/v) FA, flow was set to 1 ml min⁻¹, the wavelength of the detector was set

to λ = 214 nm and the injected volume of the sample was 20 μ L. The gradient was increased in 8 min from 5 to 90% B, then held at 90% B for 5 min, reduced to 5% B within 1 min, and held at 5% B for 4 min. Detection was done at λ = 214 nm. MS-detection range was set to 60 to 1000 (*m/z*).

MMP activation

Pro-Matrix-metalloproteinases (MMP-1, MMP-8, and MMP-9) (Merck Millipore, Burlington, MA, USA) were activated prior usage. To activate Pro-MMPs a fresh 50 mM 4-aminophenylmercuric acetate (APMA) stock solution in 0.1 M NaOH was prepared. The stock solution was added in a 10:1 (v/v) ratio to Pro-MMPs (Pro-MMPs:APMA) to reach a final concentration of 5 mM APMA. The solution was incubated for 3 h at 37 °C without agitation. Following incubation, the activated MMPs were stored in aliquots at -80 °C.

Peptide cleavage

Lyophilized peptides were solubilized in MMP buffer (200 mM NaCl, 50 mM Tris-HCl, 5 mM CaCl₂, 1 μ M ZnCl₂, 0.05% Brij35 at pH 6.8–7.0) to 1 mM stock solutions. Cleavage was performed at 37 °C under rigorous shaking in 30 μ L MMP buffer containing 0.1 mM of the respective peptide and 900 ng mL⁻¹ MMP-8. The reaction was stopped by addition of 5 mM EDTA final concentration. Iterative improvement and sensor stage experiments were performed *n* = 3, and experiments comparing Y1–8 and X3 were performed *n* = 5.

Analytical HPLC was carried out *via* an Agilent 1260 infinity II HPLC (Agilent Technologies Inc., Waldbronn, Germany) using a Zorbax 300SB-CN 5 μ m analytical 4.6 \times 150 mm LC column (Agilent Technologies Inc., Waldbronn, Germany) The device was equipped with a diode array detector (G7115A, Agilent), a fluorescence detector (G7121A, Agilent), an automatic vial sampler (G7129C, Agilent), flexible pump (G7104C, Agilent), and multicolumn oven (G7116A, Agilent). Mobile phase A was 0.1% TFA in Millipore water. Mobile phase B was ACN with 0.1% TFA, flow was set to 1 ml min⁻¹, injection volume was 15 μ L, and the wavelength of the detector was set to λ = 214 nm for PICS derived peptides.

The gradient was held at 15% B for 1 minute, then increasing to 40% within 7 minutes, increased again for 0.5 minutes to 95% B, held for 0.5 min, then back to 15% B within 0.5 minute, and held for 5.5 minutes.

Cleavage percent was analyzed by comparing the area under the curve (AUC) of the main peak from the compound incubated with MMP and in absence of MMP.

Sensor synthesis

Denatonium labeled sensors were synthesized *via* coupling of purified, side-chain-protected peptides with 2 molar equivalents of denatonium-CH₂-NH₂ and 4 molar equivalents of DIPEA in DMF. The reaction was performed at RT for 16 h under rigorous shaking. The sensors were then precipitated with DEE as described above.



DBCO-functionalized dyes were incubated with 1.5 molar excess of the respective azide-functionalized peptides in 50/50 DMSO/water (v/v) for 16 h under shaking at room temperature. The sensors were then purified by reverse phase purification as described before.

Alkyne-functionalized dyes were incubated with 1.5 molar excess of the respective azide-functionalized peptides in water with 500 μM tris-hydroxypropyltriazolylmethylamine, 100 μM CuSO_4 and 5 mM sodium ascorbate. The reaction was incubated for 1 h at room temperature and stopped by adding 500 μM

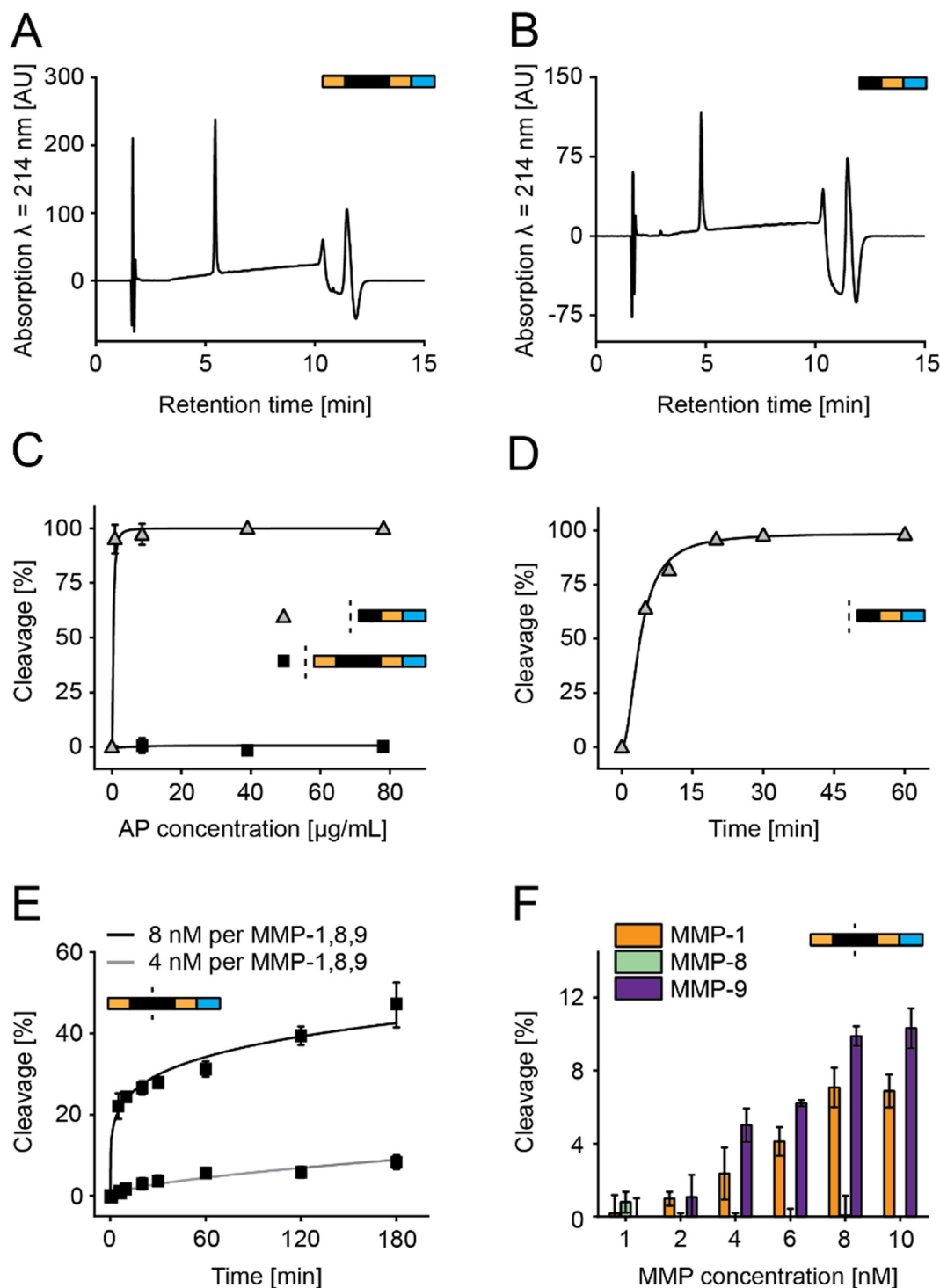


Fig. 2 (A) High performance liquid chromatography (HPLC) chromatogram of the sensor (S1). (B) HPLC chromatogram of the fragment having the bitter substance of the sensor (S1c) after MMP cleavage. (C) Aminopeptidase (AP) cleavage of S1 and S1c at different AP concentrations. (D) AP cleavage of S1c over time. (E) MMP cleavage of S1 over time with a mixture of MMP-1, 8, 9. (F) S1 cleavage following exposure to MMP-1, 8, or 9 for 15 minutes.



ethylenediaminetetraacetic acid (EDTA). The sensors were then purified *via* anion exchange chromatography *via* fast protein liquid chromatography on a GE ÄKTA pure (GE Healthcare, Chalfont St Giles, UK) system with a Hitrap Q FF (GE Healthcare, Chalfont St Giles, UK) column, with eluant A (20 mM Tris HCL pH 8) and eluent B (20 mM Tris HCL pH 8, 1 M NaCl) from 0% B to 100% B in 60 minutes. Afterwards, the sensor was purified *via* reverse phase purification as described before.

Sensors were freeze-dried (as described above) and stored at -80°C until further usage. Subsequently, sensors were characterized by HPLC and LC-MS.

Sensor cleavage

Lyophilized sensors and Y3 were solubilized in MMP buffer (200 mM NaCl, 50 mM Tris-HCl, 5 mM CaCl_2 , 1 μM ZnCl_2 , 0.05% Brij35 at pH 6.8–7.0) to 1 mM stock solutions. Cleavage was performed at 37°C under rigorous shaking in 30 μL MMP buffer containing 0.1 mM of the respective sensor/peptide and 900 ng mL^{-1} MMP-8, or a molar equivalent of MMP-1 or MMP-9 for single cleavage conditions. Cleavage of sensors in mixture was performed with sensor concentrations of 25 μM for each sensor. The

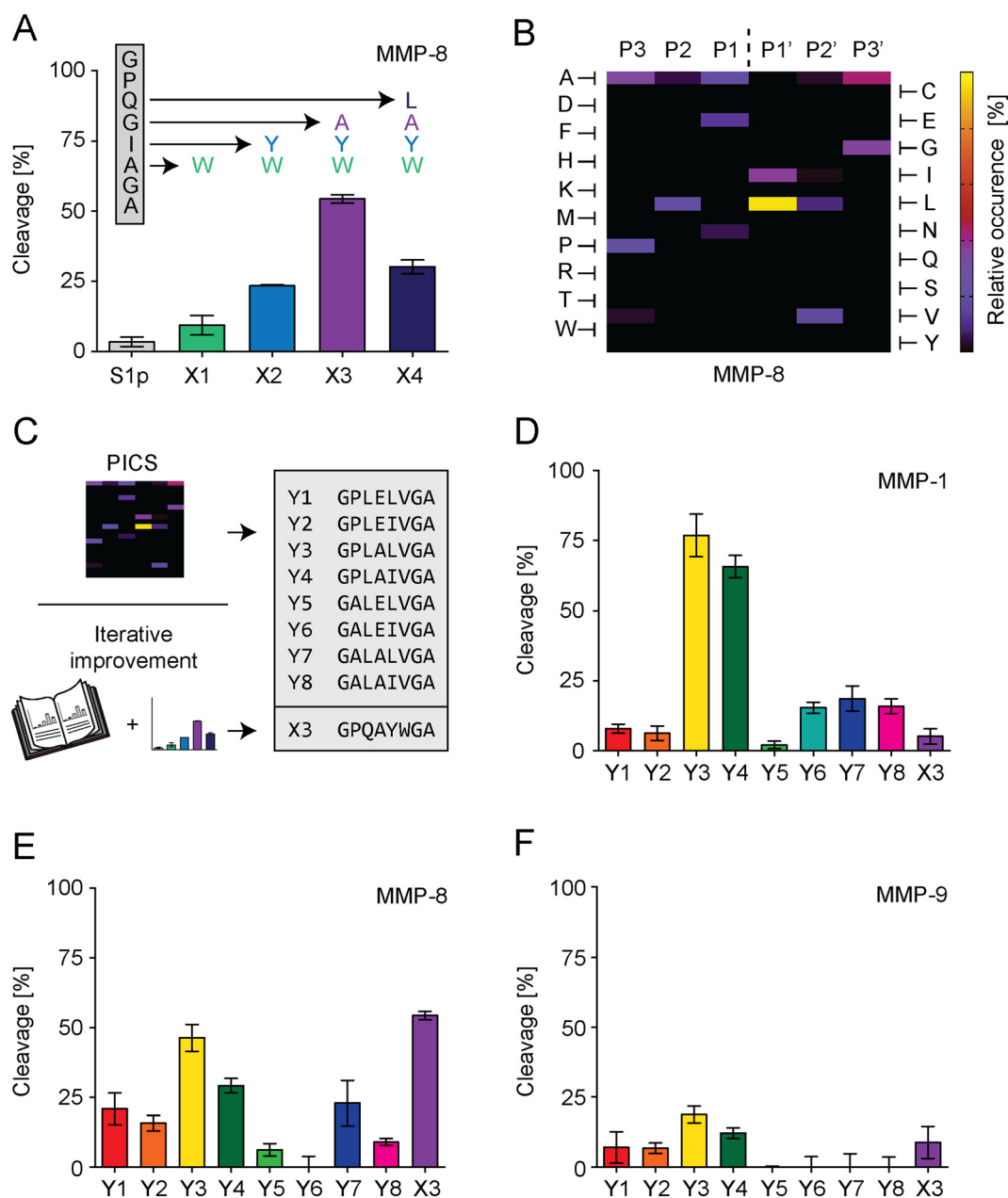


Fig. 3 (A) Cleavage of protease-sensitive sequences (PSS) derived from S1 (GPQGIAGA). (B) PICS derived relative occurrence of amino acids for position P3 to P3' using the *E. coli* proteome. (C) The X series of new sequences is derived from optimization of S1 based on known relations of primary sequence and MMP-8 sensitivity (as in panel A). The Y series is derived from our proteome searches (as in panel B). (D) MMP-1 cleavage, (E) MMP-8 cleavage, and (F) MMP-9 cleavage for selected PSSs.



reaction was stopped *via* addition of 5 mM EDTA final concentration. Experiments were performed $n = 5$. Cleavage was analyzed *via* fluorescent detection *via* HPLC with following wavelengths: SP3 (absorption: 646 nm; emission: 661 nm), SP5 (absorption: 494 nm; emission: 517 nm), SP7 (absorption: 553 nm; emission: 566 nm) and SN3 (absorption: 280 nm; emission: 350 nm). Y3 cleavage was analyzed as described before. Cleavage percent was analyzed by comparing the AUC of the main peak from the compound incubated with MMP and in absence of MMP. Cleavage percent was then normalized by multiplying cleavage results with a correction factor. The correction factor was determined individually for every experiment/MMP by dividing Y3 cleavage results acquired in protease sensitive linker identification study by cleavage results of Y3 in individual sensor cleavage experiments (Fig. 3D–F).

Sensor cleavage in human saliva

Lyophilized sensors were solubilized in MMP buffer to 0.05 mM each and mixed with saliva 1:1 (v/v) to reach a final concentration of 25 μM per sensor. The solution was incubated for 15 min at 37 °C under rigorous shaking and the reaction was stopped *via* addition of 5 mM EDTA final concentration. The samples were diluted 1:2 with acetonitrile, 1:5 with water (1:10 total dilution) and analyzed *via* HPLC as described before. Cleavage efficiency was analyzed by comparing the AUC of the newly formed peaks compared to the total AUC of the combined peaks. Peaks with LU < 1 were considered below the limit of detection and set to 0.5 LU for further analysis.

Aminopeptidase cleavage

Lyophilized sensors were solubilized in AP buffer (50 mM Tris-HCl, 1 mM CaCl_2 , 150 mM NaCl at pH 7.0) to 1 mM stock solutions. Cleavage was performed at 37 °C under rigorous shaking in 30 μL AP buffer containing 0.1 mM of the respective sensor and 0, 0.87, 8.7, 39 or 78 $\mu\text{g mL}^{-1}$ AP for 1 h for concentration-based cleavage. For time resolved cleavage of S1c, cleavage was performed at 37 °C under rigorous shaking in 30 μL AP buffer containing 0.1 mM of the respective sensor and 0.87 $\mu\text{g mL}^{-1}$ AP for 0, 5, 10, 20, 30 and 60 min. The reaction was stopped *via* 15 min incubation at 95 °C and analyzed by HPLC described before. Cleavage was calculated as the decrease of the main peak in relation to a negative control, indicating cleavage as the cleavage of one or more amino acids from the N-terminus. All experiments were performed $n = 3$.

Active MMP-8 concentration determination

Saliva samples were thawed at 4 °C, overnight, diluted 1:200 and analyzed by MMP-8 activity assay (Quickzyme, Leiden, Netherlands) following provided protocols. In brief, reagents were prepared as described by protocols. A 96 well plate was prepared: anti-MMP-8 antibody solution was added, allowed to bind at 37 °C for 2 h, followed by 4 wash steps, incubation

with assay buffer and then addition of either samples (100 μL of diluted saliva) or standards as supplied. The plate was covered and incubated at 4 °C overnight and washed 4 times with wash buffer. 1.5 mM APMA solution was freshly prepared and added to wells containing standards. Samples were treated with assay buffer to measure endogenous MMP-8 levels. The plate was covered and incubated at 37 °C for 1 h, detection reagent freshly prepared and added. Absorbance was measured at 405 nm in a microplate reader (Tecan Infinite F50, Crailsheim, Germany) at 0, 2, 4 and 6 h. Between measurements, the plate was incubated at 37 °C.

Statistical analysis

Comparison between two groups and correlations were performed using MEDAS (C. Grund, Margetshöchheim, Germany). For comparisons between two groups, Student's *t*-test was performed when Gaussian distribution was provided and Mann–Whitney U test was used when data was not normally distributed. Correlations were described by analysis of Spearman's rank correlation coefficient (ρ/rho) when data had no ties and not normally distributed. When data had ties and not normally distributed, Kendall rank correlation (τ/tau) coefficient was used to describe correlations. *P*-Values < 0.05 were considered significant and displayed as * for values < 0.05, ** for values < 0.005 and *** for values < 0.001.

Results

In this study, we optimized peptide-based sensors, which release a bitter substance (denatonium derivative) after MMP-specific cleavage (Fig. 1A).^{21,35,36} This design simplified previous designs using beads and removed previous challenges on salivary stability and storage stability by introducing acetylated N-termini and a storage stable C terminal denatonium modification.^{21,36} We optimized sensors *in vitro* using MMP-8, a key surrogate marker in periodontitis (Fig. 1B), but selected the final sensors from responses observed in a multiplex based MMP-screening in saliva samples collected from diagnosed patients (Fig. 1C).

To allow the C-terminal coupling of denatonium, an amine-functionalized denatonium derivative ($\text{Den-CH}_2\text{-NH}_2$) was synthesized and characterized (Fig. S1†).³⁵ Sensor S1 and a cut variant (S1c, representing the C-terminal fragment after MMP cleavage) were synthesized and characterized by HPLC and LC-MS, showing purity of >95% and the predicted molecular mass, as well as long term storage stability (Fig. 2A and B and S2A–C†).

Aminopeptidase stability of S1 and S1c was studied for 1 h with aminopeptidase concentrations ranging from 0 to 78 $\mu\text{g mL}^{-1}$, showing no cleavage for all tested concentrations for S1 and a cleavage rate of 95% at the lowest concentration tested for S1c (Fig. 2C). Cleavage over time was evaluated for S1c with 0.87 $\mu\text{g mL}^{-1}$ aminopeptidase over 1 h, indicating a cleavage rate of 95% after 20 min (Fig. 2D). This confirmed that unspecific aminopeptidase digest of the sensor occurs



only after the specific cleavage event and finally leads to the release of the bitter substance.

S1 was then incubated with either 4 nM or 8 nM MMP mix (a mixture of MMPs 1, 8, and 9), resulting in cleavage rates of <1% and >20% after 5 min, respectively (Fig. 2E). To test the cleavage of individual MMPs, the sensor was incubated with MMP-1, MMP-8, and MMP-9 for 15 min with concentrations ranging from 1 to 10 nM (Fig. 2F). Cleavage was observed for MMP-1 and MMP-9 but not for MMP-8, with a maximum cleavage of 7% for MMP-1 and 10% for MMP-9. Therefore, we had to redesign the PSS to address the low overall MMP sensitivity and the lack of MMP-8 susceptibility.²¹

For S1 improvement, two different strategies were followed to optimize the PSS of S1 (Fig. 3C). The first strategy exchanged individual amino acids of the S1-PSS in an iterative process based on previously published information reporting increased MMP-8 specificity (Fig. 3A).³⁷ These reports rated tryptophan in position P2' as the most effective means of increasing MMP-8 cleavage rate and suggested placing tyrosine in P1', alanine in P1, and leucine in P2. Consequently, we replaced alanine in the S1-PSS with tryptophan (referred to as PSS-X1) and observed an increase in cleavage. PSS-X1 was further improved by introducing tyrosine (PSS-X2), and the PSS-X2 was further enhanced by introducing alanine (PSS-X3).³⁷ Leucine introduction, resulting in PSS-X4, reduced cleavage compared to PSS-X3. Therefore, PSS-X3 was selected. Our second strategy searched alternative PSSs by proteomic identification of protease cleavage sites (PICS; Fig. 3B and S6–S8†).³⁸ We generated chemically modified proteome libraries from *E. coli*, ¹⁵N labeled *E. coli* and HEK-293 cells. These libraries were incubated with MMP-8, and resulting cleavage products were isolated, enriched, and analyzed by MS. PICS indicated that P3 should be an alanine or proline, P2 a leucine, P1 a glutamic acid or alanine, P1' a leucine or isoleucine, P2' a valine, and P3' an alanine or glycine. These amino acids were selected by considering their relative occurrence, as determined through WebPICS.³⁹ We set the relative occurrence threshold at 11.5% or more (average of at least three independent PICS results) to limit the number of possible candidates for peptide design.³⁸ We chemically synthesized eight PSSs based on these insights, excluding the P3' glycine variant to avoid structural repetitions. Furthermore, none of the hits exceeded the threshold of 11.5% for P4 or P4', so we selected glycine and alanine for these positions, respectively. In addition, glycine and alanine are in positions P4 and P4' in the S1-PSS (Table S1†). The PICS-derived PSSs are called PSS-Y1 to PSS-Y8 (Fig. 3C), were characterized, and tested in comparison to PSS-X3 for cleavage rates at 17.5 nM MMP-1, MMP-8, or MMP-9 with an incubation time of 15 minutes (Fig. S9 and S10† and 3D–F). We selected three PSSs for further testing. PSS-Y3 was chosen for its high cleavage rate by all MMPs. PSS-Y5 and PSS-Y7 were selected based on cleavage by MMP-8 while not being cleaved by MMP-9. PSS-X3 (see above; optimized from

previously reported insights³⁷) was chosen due to its high cleavage rate for MMP-8 and low cleavage rate for MMP-1 and MMP-9.

These four selected PSSs were then assembled into fully functional sensors using four different fluorophores and referred to as SY3, SY5, SY7, and SX3 in reference to PSS-Y3, PSS-Y5, PSS-Y7, and PSS-X3 (*vide supra*). SY3, SY5, and SY7 were synthesized with C-terminal L-azidohomoalanine (Aha) and “clicked” to different fluorescent dyes at their respective C-termini. SX3 was modified with the (fluorescent) denatonium derivative. The fluorescence spectra did not overlap. Therefore, we could simultaneously multiplex all sensors under identical conditions (Fig. 4A–C and S11–S13†). Multiplexing was tested against MMP-1, MMP-8, or MMP-9 (Fig. S15D–F†, 17.5 nM; incubation 15 minutes). The cleavage outcome of the sensors generally confirmed the effect observed for the PSS alone (Fig. 3D–F). SY3 was cleaved most effectively by MMP-1 and MMP-8 and, to a lesser extent, MMP-9. SY5 responded to MMP-8 but not MMP-1 or MMP-9, with generally low cleavage rates. SY7 had a low general cleavage rate and responded to all MMPs other than MMP-9. SX3 showed high cleavage rates for MMP-8, low cleavage rates for MMP-1, and medium cleavage rates for MMP-9. Further modifications of the fluorophore distant from the PSS did not impact cleavage performances (*e.g.*, PSS, spacer-PSS-spacer, or spacer-PSS-spacer-fluorophore, Fig. S3, and S14†).

These sensors were profiled in saliva samples from 16 healthy subjects and 50 established periodontitis patients. The saliva was characterized for total MMP-8 and active MMP-8 and correlated with cleavage rates of the sensors (Fig. 5 and S16, Tables S2–S4†). We collected further patient information, including age, smoking status, BoP, the number of missing teeth, recession, plaque, tooth mobility, furcation involvement, gingival pocket depth, clinical attachment level, periodontal epithelial surface area (PESA), and periodontal inflamed surface area (PISA) separately. The selected cohort consisted of treated and stable patients, and for example a chewing gum as a future medical product could be profiled in this group for follow-up checks at home. We chose these patients as we wanted to establish a saliva library with different MMP-8 profiles and test our sensors on that library accordingly. Sensor cleavage significantly correlated with total and active MMP-8 concentration (Fig. 5; Table S4†). However, another arguably more relevant patient pool than the one used here (to establish a proof of mechanism) are naïve periodontitis patients to facilitate diagnosis in early stages. Future studies should address this patient pool. Furthermore, all sensors except SY7 were significantly better cleaved in saliva from patients compared to saliva from healthy donors (Table S5†). Lastly, gingival pocket depths significantly correlated with sensor cleavage, except for SY7 (Table S6†).

Discussion

The peptide-based sensors were stable against unspecific salivary aminopeptidases. Following MMP-cleavage of their



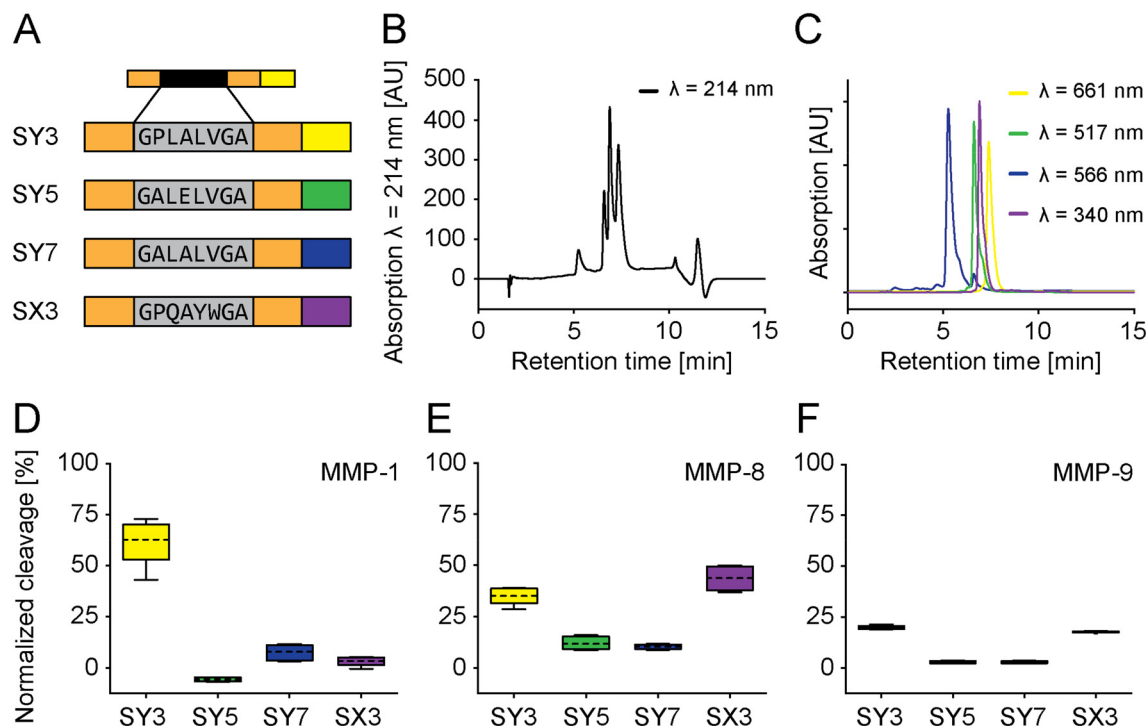


Fig. 4 (A) Sensor overview. (B) HPLC chromatogram of mixture of SY3, SY5, SY7, and SX3. (C) HPLC chromatogram acquired at individual wavelengths for a mixture of SY3, SY5, SY7, and SX3. (D) Normalized MMP-1 cleavage and (E) MMP-8 cleavage and (F) MMP-9 cleavage for SY3, SY5, SY7, and SX3. Box plots show the mean as dotted line, whiskers indicate outliers and the range indicates the 25% and 75% percentiles, respectively.

PSS, they rapidly disintegrated, releasing a bitter flavor (Fig. 2C and D). The new soluble design replaces previous particle-based sensors.^{21,36} One challenge during replacement was that the PSS, used in the previous particle-based design, was unresponsive to MMP-8 when integrated into the new and soluble sensors. Therefore, we initiated a search for novel MMP-8 responsive PSSs compatible with the new and soluble setup (Fig. 3). Identified hits (PSS) were assessed, individually encoded with fluorescent dyes (sensor) and multiplexed in patient and control saliva. Sensor cleavage correlated with MMP-8 (total and active) levels *in vitro* and MMP-8 levels measured in patient saliva (Fig. 5). Furthermore, cleavage rates were significantly higher in patient saliva compared to saliva from healthy donors (Table S5†). Further adaptations became necessary. For example, the particle in the old design protected the N-terminus of the sensor from unspecific cleavage by salivary aminopeptidases. The same was achieved in the new soluble sensor by N-terminal acetylation. As a result, replacing the old, particle-based design will reduce the costs of goods for this high volume, commodity product, facilitates manufacturing, including the integration into the development of new oral thin films or chewing gums (for example), and reduces the effort for quality control reflecting the soluble state of the new sensors.

Contrasting previous studies linking salivary MMP-8 to periodontitis, our patient pool did not show these correlations (Fig. S18, Table S5†).⁴⁰ These differences arguably root in our inclusion/exclusion criteria, recruiting treated periodontitis

patients in regular supportive aftercare with comparatively low BoP-values. Treatment reduces acute inflammatory processes and MMP levels.^{41,42} Furthermore, our criteria did not gate salivary collection regimens, such as the time of the day or last food/drink intake. MMP-8 is not the only marker correlating with periodontitis.^{28–31} Therefore, the possible promiscuity of our sensors might be advantageous, which is why we screened in patient saliva to reflect the complexity in periodontitis. Some sensors which were not exceptionally responsive to MMP-8 (SY3, SY5, and SX3) discriminated against periodontitis, supporting final screenings in biological fluids. Our patient pool donated saliva with a relevant range of active inflammation, but this patient pool is not the one targeted by the product. The future target profile is the broad, un-diagnosed population to early segment subjects at risk of developing periodontitis. The clinical correlation demonstrated further exciting potential. For example, sensor cleavage correlated with pocket probing depth, an important factor for disease progression.⁴³ If corroborated by future clinical studies using pocket depth as the primary endpoint, this evidence further strengthens the approach for community screening, targeted dental diagnosis and dental care in prescreened and enriched patient pools.

MMP-8 is a significant biomarker for a spectrum of physiological processes ranging from inflammation, wound healing, and immune responses to malignant tissue and tissue remodeling, highlighting its potential as a versatile diagnostic and therapeutic target.⁴⁴ As a diagnostic marker, MMP-8 is already used in various rapid test systems in the



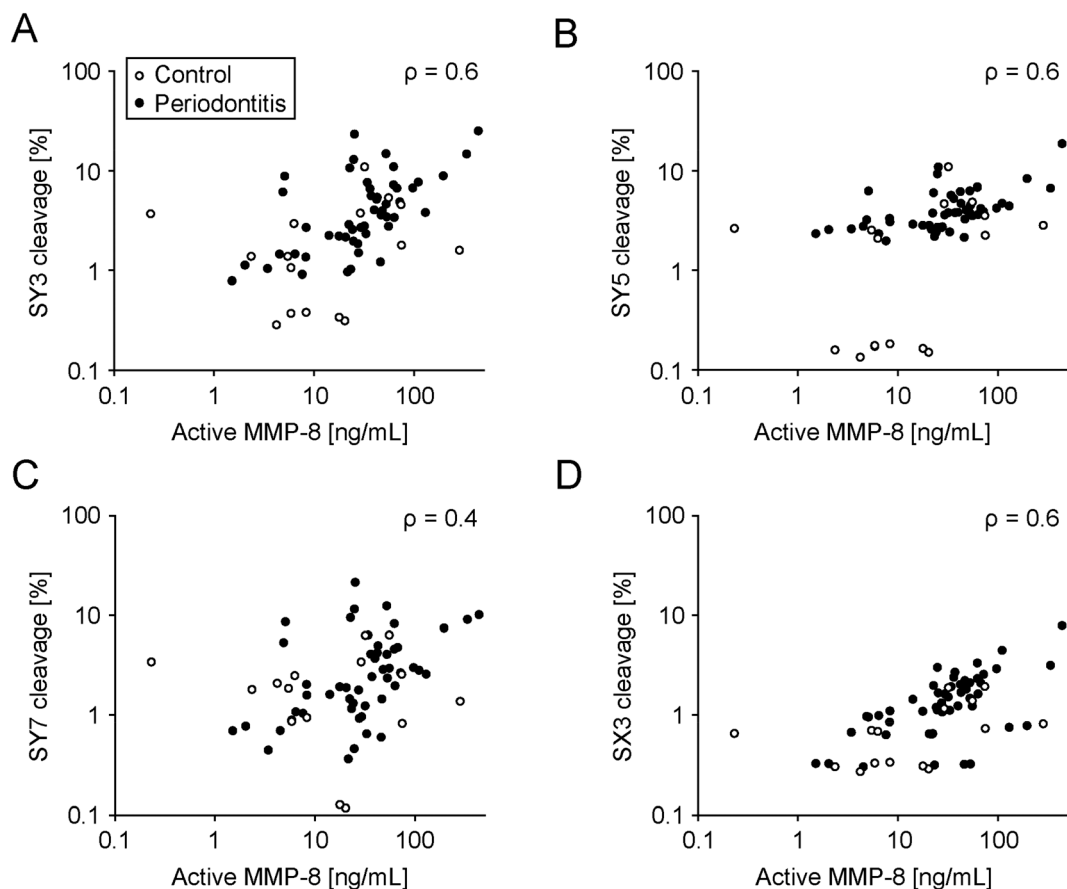


Fig. 5 Cleavage of individual sensors in human saliva. Open circles result from the control group and filled circles from the periodontitis group. (A) SY3 cleavage ($\rho = 0.6$), (B) SY5 cleavage ($\rho = 0.6$), and (C) SY7 cleavage ($\rho = 0.4$), and (D) SX3 cleavage ($\rho = 0.6$) in response to the measured active MMP-8 concentration in saliva.

field of periodontitis (*vide supra*). Our sensor does not offer the possibility of quantifying periodontitis but allows a simple, fast and ubiquitous use without a professional background. Moreover, the sensors are modular. The PSS is readily replaced by other PSSs, paving the way for new products, for example, marking bacterial infection (strep throat) or viral challenge (flu).^{45,46} We also currently explore use in spotting malignancies like squamous cell carcinomas of the head and neck.⁴⁷ The rapid PICS-identification tools for new PSSs leverage modular flexibilities and, when combined with multiplexing provide paired, head-to-head comparison within each biological sample. Nevertheless, there is bias in our studies. We rely on previous information, such as the link between MMP-8 and periodontitis.^{22,23} However, MMP-8 levels correlate with periodontitis but not necessarily *vice versa*.²³ Consequently, our approach is burdened by the limitations associated with MMP-8 as a biomarker for periodontitis.

In summary, this platform technology holds the potential to leverage active community screening for various challenges. Future designs may target viral outbreaks. The modular design and the rapid and proteome-wide search tools for new PSSs suggest quick response rates, easy mass-production, and distribution at affordable costs.

Conflicts of interest

L. M. is an inventor of related patents/patent applications (EP13710333.9, EP17758824.1, EP17762078.8, and EP19708959.2A) and states a possible conflict of interest. M. R. is an inventor of a related patent application (EP19708959.2A). The remaining authors declare no competing interests.

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