Have you had bleeding from your gums? Self-report to identify gingival inflammation (The SING diagnostic accuracy and diagnostic model development study)

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Abstract

Aim: To assess the diagnostic performance of self-reported oral health questions and develop a diagnostic model with additional risk factors to predict clinical gingival inflammation in systemically healthy adults in the United Kingdom.

Methods: Gingival inflammation was measured by trained staff and defined as bleeding on probing (present if bleeding sites ≥ 30%). Sensitivity and specificity of self-reported questions were calculated; a diagnostic model to predict gingival inflammation was developed and its performance (calibration and discrimination) assessed.

Results: We included 2853 participants. Self-reported questions about bleeding gums had the best performance: the highest sensitivity was 0.73 (95% CI 0.70, 0.75) for a Likert item and the highest specificity 0.89 (95% CI 0.87, 0.90) for a binary question. The final diagnostic model included self-reported bleeding, oral health behaviour, smoking status, previous scale and polish received. Its area under the curve was 0.65 (95% CI 0.63–0.67).

Conclusion: This is the largest assessment of diagnostic performance of self-reported oral health questions and the first diagnostic model developed to diagnose gingival inflammation. A self-reported bleeding gums question or our model could be used to rule in gingival inflammation since they showed good sensitivity, but are limited in identifying healthy individuals and should be externally validated.

Keywords
diagnosis, epidemiology, gingival inflammation, prediction modelling, self-report

Clinical Relevance

Scientific rationale for Study: Gingival inflammation is a precursor of periodontitis and needs to be identified in population-based studies; however, it is costly and difficult to measure. Self-reported questions can be an efficient and inexpensive alternative to clinically assessed measures in epidemiological surveillance.

Principal Findings: A self-reported bleeding gums question had high sensitivity and low specificity to identify clinically measured gingival inflammation. Our diagnostic prediction model for gingival inflammation had moderate discriminatory ability.
1 | INTRODUCTION

Periodontal disease is an inflammatory disease that affects the soft and hard tissues supporting teeth. This disease is largely preventable, yet it remains the major cause of poor oral health worldwide and is the primary cause of tooth loss in older adults (Petersen & Ogawa, 2000). Periodontal disease is classified into two broad categories: (Petersen & Ogawa, 2000) gingivitis and (Tonetti et al., 2020) periodontitis. Gingivitis is usually characterized by, amongst other factors, gingival inflammation, which is a reversible condition identified by bleeding at the gingival margin. Gingival inflammation is an early sign of periodontal disease, (Tonetti et al., 2020) and its association with periodontal health has been explored in numerous publications (Kallio, 1996; Weintraub et al., 2013). Identifying gingival inflammation could help prevent periodontal disease.

The development of self-reported tools to measure periodontal disease risk is particularly important in the field of oral health surveillance (Ramos et al., 2013), since clinical measures are difficult and costly to collect, and hard or impossible to standardize. The Centres for Disease Control and Prevention (CDC) and the American Academy of Periodontology recommended since 2003 the use of self-reported measurements that could be valid to predict the prevalence of periodontal disease as an alternative to examinations (Eke et al., 2012). Several self-reported oral health questions have been previously discussed as an alternative to clinical measures of periodontal disease (Blicher et al., 2015; Abbood et al., 2016), but studies focusing on gingivitis or gingival inflammation have been inconclusive due to small, non-representative samples and the use of single questions without the consideration of other potential risk factors (Abbood et al., 2016).

Prediction models and specifically diagnostic models allow the inclusion of several risk factors to predict the existence of a condition (Collins et al., 2015). Prediction models are currently used in different areas of medicine (Collins et al., 2015) and have been increasingly popular in assessing the prevalence and progression of periodontitis (Du et al., 2018). There are currently no models available to predict gingival inflammation. Diagnostic models of gingival inflammation could be used as a first line assessment in a clinical surveillance system, identifying patients that need further assessment, or they could be incorporated in large epidemiological studies to target clinical examinations in research participants.

We aimed to assess the diagnostic accuracy of several self-reported oral health questions individually to identify gingival inflammation, measured by bleeding on probing, in a large and representative UK-based sample. We developed and validated a diagnostic model including other gingival inflammation risk factors to assess whether additional risk factors would improve the diagnostic accuracy of these measures. We hypothesized that using self-reported oral health questions with or without additional risk factors could help identify clinical gingival inflammation and, therefore, constitute an alternative to clinical measurement in research settings.

2 | METHODS

2.1 | Data

We collected and combined clinical and self-reported data from adults participating in two large, UK-wide dental randomized trials. Combination of data from both trials was deemed reasonable to follow TRIPOD guidelines and maximize statistical precision (Collins et al., 2015) and because inclusion criteria, setting, recruitment and data collection processes, patient reported and clinical outcomes collected were identical in both trials. Moreover, the trials were conducted by the same team of researchers.

2.1.1 | IQuaD trial

The IQuaD trial (Ramsay et al., 2018) compared the clinical and cost-effectiveness of providing scale and polish 6 monthly, 12 monthly or never during 3 years; the clinical and cost-effectiveness of personalized vs standard oral hygiene advice were also compared. The study recruited 1877 participants from Scotland and Northeast England. Recruitment began in 2011, and data were collected until September 2016.

2.1.2 | INTERVAL trial

The INTERVAL trial (Clarkson et al., 2018) compared the clinical and cost-effectiveness of 24-monthly recalls vs 6-monthly vs risk-based on the same outcome and randomized 2372 participants from Scotland, England, Wales and Northern Ireland. Recruitment began in July 2010, and data were collected until August 2018.

Both trials were pre-registered and had bleeding on probing as their primary outcome. They were set up in primary dental care provided by the National Health System in the United Kingdom. Participants were recruited via their dental practices.

Participant-reported data, including the index tests, in IQuaD and INTERVAL were collected using an annual patient questionnaire from baseline until year 3 (for IQuaD) or 4 (for INTERVAL). The data used in our model are restricted to the final year follow-up of
TABLE 1 Four self-reported gingivitis measures used and their scores and transformation for individual diagnostic analysis

| Self-reported measures                              | Original score (used in the diagnostic model)                              | Transformation for individual diagnostic accuracy analysis |
|-----------------------------------------------------|--------------------------------------------------------------------------|------------------------------------------------------------|
| Self-report bleeding gums (Likert)                  | 1:Never; 2:Hardly ever; 3:Occasionally; 4:Fairly often; 5:Very often.     | A score of 2 or more was considered as a positive index test |
| (‘Have you had bleeding from your gums when brushing your teeth?’) |                                                                          |                                                             |
| Self-report bleeding gums (Binary)                  | Yes/No                                                                   |                                                             |
| (‘Do your gums bleed when brushing?’)              |                                                                          |                                                             |
| Self-report unpleasant taste in your mouth          | 1:Never; 2:Hardly ever; 3:Occasionally; 4:Fairly often; 5:Very often.     | A score of 2 or more was considered as a positive index test |
| (‘Have you had an unpleasant taste in your mouth when brushing your teeth?’) |                                                                          |                                                             |
| Self-report bad breath                              | 1:Never; 2:Hardly ever; 3:Occasionally; 4:Fairly often; 5:Very often.     | A score of 2 or more was considered as a positive index test |
| (‘Have you had bad breath?’)                       |                                                                          |                                                             |

each trial; therefore, this is a cross-sectional analysis. IQuaD and INTERVAL collected similar information about the participants, but not all outcomes were collected in both trials.

Clinical data were collected at follow-up (3 or 4 years post-randomization for IQuaD and INTERVAL, respectively). Variables collected at the clinical assessment included the following: bleeding on probing, calculus and pocket depth. Clinical outcomes were measured at the end of each trial by trained outcome assessors. Gingival inflammation was measured according to the Gingival Index of Loe (1967) by running the UNC probe circumferentially around each tooth just within the gingival sulcus or pocket. After 30 s, bleeding was recorded as being present or absent on the buccal and lingual surfaces. Percentage of sites bleeding on probing per participant was calculated by adding the sites where bleeding was present in each participant (two sites per tooth – buccal and lingual) and dividing by twice the number of teeth in the mouth and then multiplied by 100, thus generating a variable varying from 0 (no bleeding in any of the sites available) to 100 (bleeding in all sites available). More information about clinical outcome collection can be found in IQuaD’s and INTERVAL’s protocols (Clarkson et al., 2013; Clarkson et al., 2018). Participants did not have access to their clinical data before answering the questions and clinical examiners did not have access to self-reported data.

2.2 Study population

Eligibility criteria were similar in both trials: both included adult participants (≥18 years of age) who were dentate, had attended a check-up at least twice in the 2 years prior to the trial and received their dental care in part or fully as a patient in the National Health System in the United Kingdom. Patients with an uncontrolled medical condition (e.g. diabetes and immunocompromised) were excluded from both trials. IQuaD had an additional inclusion criterion to INTERVAL: participants had to score 0–3 in their Basic Periodontal Examination. Participants were approached and recruited if they met the eligibility criteria and accepted to take part; therefore, we used a convenience sample.

2.3 Gingival inflammation definition

According to the most recent classification from the 2017 Classification of Periodontal Diseases (Chapple et al., 2018; Trombelli et al., 2018), for adults with pocket depths ≤3 mm, and an intact and a reduced periodontium, generalized gingivitis for epidemiological purposes can be defined as 30% or more bleeding sites. Localized gingivitis is defined by having between 10 and 29% of bleeding sites. Participants with <10% sites bleeding are gingivitis-free. Because pocket depths could be higher than 3 mm and we did not measure clinical attachment level, we did not focus on gingivitis. We used 30% as the primary cut-off for our analyses and considered it to be an indicator of gingival inflammation. To explore the impact of lowering the threshold to define gingival inflammation in our models’ secondary analyses used 10% sites bleeding as the cut-off.

2.4 Diagnostic accuracy of individual self-reported oral health questions

2.4.1 Outcome

Outcomes are the sensitivity and specificity of the index tests on detection of gingival inflammation (defined as having bleeding on probing equal or higher than 30%). We explored the impact of lowering the threshold to define gingival inflammation (using 10% or greater sites bleeding as the definition) in a sensitivity analysis.

2.4.2 Index tests

Four self-reported bleeding or bleeding-related measures were used as the index tests in the current study. Table 1 presents the measures and their score. In order to perform the diagnostic accuracy analysis for single self-reported gingival inflammation measures, Likert items were transformed into binary measures. To decide the appropriate threshold, we assessed the scales’ diagnostic performance at.
different thresholds using receiver operating characteristics curves and selected the best performance. We assumed the best performance to be the one that had the overall best proportion of correctly classified individuals considering both sensitivity and specificity.

2.4.3 | Statistical analysis methods

Diagnostic measures (sensitivity and specificity) were calculated for each self-reported index test measure with 95% confidence intervals calculated using the Agresti–Coul method (Newcombe, 1998). We used a complete case approach where diagnostic measures were calculated if participants had information in both the index test and reference standard (clinical gingival inflammation).

2.5 | Diagnostic model to predict gingival inflammation

2.5.1 | Outcome

The main outcome of interest was generalized gingival inflammation defined as 30% or more sites with bleeding on probing. We explored the impact of lowering the threshold to define gingival inflammation (using 10% or greater sites bleeding as the definition) in a sensitivity analysis.

2.5.2 | Candidate predictors

All variables collected in IQuaD and INTERVAL were considered for inclusion in the model (Table S1). A group of clinical experts, selected by convenience, was approached initially via email and asked to select the most relevant predictors of gingival inflammation. In a face-to-face meeting, the experts were reminded of the survey and there was an opportunity to discuss the topic in smaller groups. Sixteen experts replied to a survey, via Google Docs, available during February 2019. To be considered for inclusion in the model, a predictor had to be selected as important by at least one expert. All predictors selected were included in the ‘full model’. The main model excluded clinical predictors (number of teeth, calculus and probing depth). As a sensitivity analysis, we included clinical predictors in the model. For the diagnostic model, self-reported gingival inflammation measures were included as originally scored (as shown in Table 1). Randomized treatment was not included in the model given the cross-sectional nature of the analysis, and the fact that any treatment was only provided to patients after the clinical outcomes was measured.

2.5.3 | Statistical analysis methods

2.6 | Development of a prediction model to identify gingival inflammation

The univariable (unadjusted) associations between each candidate predictor and gingival inflammation were assessed using a logistic regression model, to assess the impact of each candidate predictor individually in relation to gingival inflammation. We assessed linearity between predictors and the log odds of gingival inflammation and found all relationships to be linear. A p-value of <.05 identified statistically significant univariate associations.

2.7 | Candidate predictor selection

All candidate predictors selected by experts were included in an initial ‘full model’. To select the final predictors in the final multiple logistic regression model, I implemented an automated backward selection using p-value <.10 as the selection criteria. I used this selection threshold because it is a common one and simulation studies show values above 0.05 should be considered (54).

2.8 | Sample size

Sample size was constrained by the number of participants in IQuaD and INTERVAL with a final year clinical assessment. We included participants who had a primary outcome (clinical measure of gingival inflammation), at least one of the self-reported index test measures and predictors collected in both trials.

2.9 | Missing data

We assumed a missing at random mechanism and used multiple imputation as a sensitivity analysis to impute missing data. We excluded predictors from the multiple logistic regression complete case main model that had more than 10% missing data.

2.10 | Performance measures

Performance was evaluated using the full dataset following the appropriate recommendations (Collins et al., 2015). We quantified the ability of the multiple logistic regression model to discriminate between participants with and without gingival inflammation (discriminatory ability) by calculating the area under the receiver operating characteristics curve (c-statistic), sensitivity and specificity (54). The c-statistic gives the probability of a participant with the condition to be given a higher probability of having the condition by the model compared with a randomly chosen participant without the condition (93). Sensitivity and specificity were calculated based on a predicted probability threshold of 0.5. We used this threshold because, at this stage of the model development, we are equally interested in detecting individuals with and without bleeding gums. Calibration of the model (i.e. the agreement between observed gingival inflammation and predicted gingival inflammation) was assessed using a calibration plot (93). The plot assessed this for each 10th
percentile of predicted risk, generating 10 equally sized groups, by plotting observed proportions versus predicted probabilities and adding a smoothed (lowess) line over the entire predicted probability range (93).

2.11 | Internal validation

To assess the model calibration’s optimism (defined as the bootstrap performance of the model minus the test performance), we used bootstrapping for internal validation. Two hundred bootstrap samples, each containing the same number of patients, were generated with replacement. The c-statistic was calculated for each of the 200 sample-derived models (bootstrap performance) and also calculated for each of the 200 sample-derived models applied to the original dataset (test performance). The difference between the two c-statistics for each sample was averaged over the 200 samples. This reflected the optimism of the original model’s performance.

Analyses were performed using Stata/SE 15 (StataCorp 2016). A study protocol is available upon request.

2.12 | Ethical considerations

Favourable ethics opinion for the IQuaD trial was confirmed by the East of Scotland Research Ethics Service on 24 March 2011 [Research Ethics Committee (REC) reference number 10/S0501/65]. Favourable ethical opinion for the INTERVAL Dental Recalls Trial was granted by the Fife and Forth Valley Research Ethics Committee [REC Reference number 09/S0501/1]. Written consent was obtained from participants, and more information on the process has been published elsewhere (Ramsay et al., 2018; Clarkson et al., 2018).

3 | RESULTS

We included a maximum of 2853 participants that provided a measure of clinical bleeding and at least one of the index tests (self-reported questions related to gingival inflammation; Figure 1). From those, 1694 (56%) participants had generalized gingival inflammation and 1328 (44%) did not. Overall, 6% of participants with a clinical measure of gingival inflammation were missing all index tests and 80% had all index tests available. The most missed index tests were the questions about unpleasant taste in mouth when brushing teeth and bad breath with around 15% of participants not providing an answer from those participants that provided a clinical measure. Participants that provided an answer to the index test questions had similar bleeding levels to those that skipped those questions (Table S2). Time between being clinical examined and answering the self-reported questions had a median of 22 days (Percentile 25: 5 days, Percentile 75: 42 days), meaning most participants had their clinical exam after replying to the questionnaire.

3.1 | Summary of candidate predictors

Table 2 shows a summary of candidate predictors. Participants were on average 51 years old, 60% were women, 15% identified as smokers, and 91% identified as regular attenders to the dentist. They had on average 6 out of 9 in an oral health behaviour scale, and they had good quality of life related to oral health (5.2 out of 56, where 56 is worst quality of life). Participants had on average 24 teeth (excludes third molars), 36% of sites bleeding, 37% of sites with calculus and a healthy mean pocket depth of 2 mm. Missing data rates for each candidate predictor are presented in the Table S3. We excluded candidate predictors with more than 10% missing data from the main model. For that reason, your last course of treatment was excluded.

3.2 | Diagnostic performance of the self-reported oral health questions

Figures S1-S3 show the ROC curves for each Likert scale question before transformation. The diagnostic performance for the four index tests is given in Table 3 using the main reference standard cut-off of 30%. The binary self-reported bleeding gums question had the highest specificity (0.89) but the lowest sensitivity (0.20). The Likert scale self-reported bleeding gums question had the highest sensitivity (0.73) but poor specificity (0.39). Self-reported unpleasant taste in mouth and bad breath performed poorly. Table S4 shows the diagnostic performance of self-reported oral health questions when using a threshold of 10% with similar results.

3.3 | Diagnostic modelling results

All predictors available were selected at least once by the experts invited and therefore included in the full model. Table S5 shows unadjusted odds ratios with 95% confidence intervals for each predictor. Older age, electric toothbrushes (vs manual), or using both electric and manual toothbrushes (vs manual only) and being a smoker were all significant and associated with lower odds of having gingival inflammation. Regarding self-reported oral health questions, all but bad breath had a significant association with clinical gingival inflammation. Higher oral health behaviour and perceived behaviour control scores were protective of gingival inflammation. Participants with higher calculus levels and deeper pocket depths had higher odds of having gingival inflammation.

In the final model (Table 4), the odds of having gingival inflammation were 1.4 times higher for every additional point in the Likert scale (1- never experienced bleeding from my gums to 5-always experience). Being a smoker and having a higher oral health behaviour score (i.e. doing more appropriate oral health behaviours) were associated with lower odds of having gingival inflammation. Each additional scale and polish in the previous year to participating in the trial were associated with higher odds of having gingival inflammation.
The final model was able to correctly identify participants with gingival inflammation (sensitivity) in 0.74 (95% CI 0.71–0.76) of the cases and able to correctly identify participants without gingival inflammation (specificity) in 0.47 (95% CI 0.44–0.50) of the cases. The trade-offs between sensitivity and specificity for the main model are shown in Figure 2 with the area under the curve. The probability that a patient with gingival inflammation is given a higher probability of having gingival inflammation by the model than a randomly chosen patient without gingival inflammation is 0.65 (c-statistic; 95% CI 0.63–0.67). Optimism in c-statistic was calculated at 0.012 suggesting no model overfitting. A good agreement between observed and predicted probabilities of gingival inflammation was observed in the calibration plot (Figure S4). Using multiple imputation resulted in a similar final model including the same predictors as the main model plus self-reported bleeding gums (binary) with a c-statistic of 0.65. Including clinical predictors resulted in a small increase in the c-statistic (from 0.65 to 0.69) with a similar sensitivity as the model without clinical predictors (0.72), but better specificity (from 0.47 to 0.54). A secondary analysis model using bleeding on probing above 10% as the primary outcome yielded a similar c-statistic of 0.66 (results available upon request). Given the models were based in the aggregation of two studies, iQuaD and INTERVAL, we provide the diagnostic performance of the models and self-reported questions for each study in Table S6 and their sensitivity/specificity trade-offs in area under the curve figures (Figures S5–S8).

Figure 3 shows the different discriminatory abilities of self-reported questions and the diagnostic model. The diagnostic model has similar sensitivity to the self-reported bleeding gums question (using a Likert scale) and a higher specificity, but both are poor (below 0.5).

4 | DISCUSSION

To our knowledge, this is the largest study evaluating the diagnostic performance of self-reported oral health questions to detect
suggesting they are more acceptable to patients. These findings focusing on gingival inflammation side-effects: they yielded better diagnostic performance and had lower rates of missing data, suggesting they are more acceptable to patients. These findings are in line with previous studies from Gilbert and Nuttall (Gilbert & Nuttall, 1999) and Dietrich (Dietrich et al., 2005). Even though halitosis has been shown to be associated with bleeding gums (Kayombo & Mumghamba, 2017), it is possible that the indirect nature of the questions or their potential negative connotation is unhelpful.

A single self-reported bleeding question (Likert item) had comparable diagnostic accuracy performance to the diagnostic model we developed with four predictors. The diagnostic model had marginally better sensitivity and better specificity than the self-reported bleeding Likert item, resulting in a higher, moderate discriminant ability (Model’s C-statistic = 0.65 vs Likert item area under the curve (AUC) = 0.60). Adding clinical predictors had a limited impact in the model’s diagnostic performance. In practice, researchers that can access the risk factors information included in our model will benefit marginally from it compared with asking patients about their bleeding gums with a single Likert item, but this benefit is unlikely to be of clinical importance.

Smoking status, oral health behaviour, previous pattern of having scale and polishes and self-reported bleeding were selected as predictors in the final diagnostic model. The association between smoking and periodontal health is well established (Abbood et al., 2016; Du et al., 2018), and previous periodontitis prediction models had found oral health behaviour and dental visits as significant predictors (Du et al., 2018).

The original studies (IQaD and INTERVAL) were not specifically designed as diagnostic accuracy or diagnostic model development studies, although we followed recommended practice where possible, including blinding assessors to index test results. Candidate predictors available were limited to those collected in IQaD and INTERVAL and probably excluded important predictors that could have improved the diagnostic accuracy of the prediction model. SING’s outcome, gingival inflammation, is challenging for two reasons: its definition and its measurement. We used gingivitis most recent threshold classification (Chapple et al., 2018) as an indicator of gingival inflammation. IQaD and INTERVAL included intensive measurement training of outcome assessors and examined two tooth surfaces per tooth (partial mouth examination). This is recognized to be highly desirable for both patients and oral health professionals, and particularly in the context of two large UK-wide trials, even though the extent to which the results would differ from the results of a full-mouth examination is unknown (Trombelli et al., 2018). Bleeding on probing is impossible to calibrate. This may result in measurement error and misclassification of inflammation status which can lead to biased risk scores and c-statistics (Zawistowski et al., 2017). Because the extent of misclassification is unknown, we could not address it in our analysis, which is a common problem in this field (Kuchenhoff et al., 2009).

The diagnostic performance results found in SING are similar to other periodontitis prediction models (Du et al., 2018) and show challenges across the periodontal health field in identifying prediction models that are clinically useful. This could be due to a non-comprehensive

### TABLE 2 Summary of candidate predictors – mean (SD), count unless indicated otherwise

| Candidate predictors | N = 2853 |
|----------------------|----------|
| **General**          |          |
| Age                  | 50.6 (14.6), 2853 |
| Female – n (%)       | 1719 (60.3) |
| Your last course of treatment was – n (%) | |
| NHS                  | 2148 (75.3) |
| Private              | 126 (4.4) |
| Combination          | 173 (6.1) |
| Self-reported regular attender – n (%) | 2504 (87.8) |
| Type of toothbrush – n (%) | |
| Manual               | 1363 (47.8) |
| Electric             | 1094 (38.3) |
| Both                 | 171 (6.0) |
| Patient has dental insurance – n (%) | 81 (2.8) |
| Smoked in the last 12 months – n (%) | 320 (11.2) |
| How difficult is for you to travel to the dentist | 6.4 (1.1), 2825 |
| No. of scale and polishes in the year prior to the trial (routine data) | 2.6 (2.3), 2688 |
| **Self-reported oral health** | |
| I had bleeding from my gums when brushing – n (%) | 453 (15.9) |
| Have you had bleeding from your gums when brushing your teeth? | 2.1 (1.0), 2749 |
| Have you had bad breath? | 1.9 (0.9), 2576 |
| Have you had an unpleasant taste in your mouth when brushing your teeth? | 1.4 (0.7), 2584 |
| Patient reported     |          |
| OHIP-14              | 5.2 (7.0), 2562 |
| Oral health behaviour score | 5.3 (1.7), 2636 |
| After brushing I spit but not rinse | 1060 (37.2) |
| Perceived behaviour control | 4.6 (1.4), 2635 |
| Attitude             | 4.8 (1.4), 2636 |
| **Clinical**         |          |
| Calculus (% sites)   | 36.9 (27.4), 2849 |
| Pocket depth (mm)    | 2.0 (0.4), 2843 |
| Number of teeth      | 23.6 (4.7), 2853 |
There is space to improve the sensitivity of the diagnostic model developed here. Researchers have suggested other potential predictors to improve sensitivity of prediction models in this field: gingival cervical fluid, salivary markers or microbiology information have been identified as useful in periodontitis models (Du et al., 2018); systemic factors such as metabolic or nutritional factors and hematologic conditions can affect the extension, severity and progression of gingivitis and gingival inflammation (Chapple et al., 2018). However, it is important to find a balance between discrimination ability and cost of a diagnostic tool, as well as ease of implementation.

| TABLE 4 | Odds ratios and 95% confidence intervals (CI) of each predictor on gingivitis adjusted for all available predictors (final model) |
|-----------|-------------------------------------------------|
| Predictors | Odds ratio 95% CI, p-value |
| Bleeding gums (Likert scale) | 1.40 (1.28, 1.53), <.001 |
| Non-smoker vs smoker (reference category) | 1.73 (1.34, 2.24), <.001 |
| Scale and polish (previous year to the trial) | 1.05 (1.01, 1.08), .02 |
| Oral health behaviour score | 0.80 (0.76, 0.84), <.001 |

There is space to improve the sensitivity of the diagnostic model developed here. Researchers have suggested other potential predictors to improve sensitivity of prediction models in this field: gingival cervical fluid, salivary markers or microbiology information have been identified as useful in periodontitis models (Du et al., 2018); systemic factors such as metabolic or nutritional factors and hematologic conditions can affect the extension, severity and progression of gingivitis and gingival inflammation (Chapple et al., 2018). However, it is important to find a balance between discrimination ability and cost of a diagnostic tool, as well as ease of implementation.
involved in planning, conducting and delivering both trials.

We would like to acknowledge all study participants in IQuaD and INTERVAL for contributing with their information, and the trial teams involved in the interpretation of results. BG drafted the manuscript. GM and CR commented on the manuscript. All authors agreed to the final version of the manuscript.

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CONFLICT OF INTEREST
The authors have stated explicitly that there are no conflicts of interest in connection with this article.

AUTHOR CONTRIBUTIONS
BG, GM and CR designed the study. CR co-led the data collection process. BG analysed the data. All authors were involved in the interpretation of results. BG drafted the manuscript. GM and CR commented on the manuscript. All authors agreed to the final version of the manuscript.

DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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