

Article **Repeatability of taste recognition threshold measurements with** *QUEST* and *quick Yes-No*

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- Abstract: Taste perception, though vital for nutrient sensing, has long been overlooked in sensory
- ² assessments. This can, at least in part, be attributed to challenges associated with the handling of
- ³ liquid, perishable stimuli, but also with scarce efforts to optimize testing procedures to be more
- time-efficient. We have previously introduced an adaptive, QUEST-based procedure to measure taste
- sensitivity thresholds that can be quicker than other existing approaches, yet similarly reliable [1].
- 6 Despite its advantages, this procedure lacks experimental control of false alarms (i.e., response bias)
- 7 and psychometric function slope. Variations of these parameters, however, may also influence the
- threshold estimate. This raises the question as to whether a procedure that simultaneously assesses
- threshold, false-alarm rate, and slope might be able to produce threshold estimates with higher
- ¹⁰ repeatability, i.e., smaller variation between repeated measurements of the same participant. Here,
- we compared the performance of QUEST with a method that allows measurement of false-alarm
- rates and slopes, *quick Yes-No* (qYN), in a test-retest design for citric acid, sodium chloride, quinine
- hydrochloride, and sucrose recognition thresholds using complementary measures of repeatability,
- namely test-retest correlations and coefficients of repeatability [2]. Both threshold procedures yielded
- largely overlapping thresholds with good repeatability between measurements. Furthermore, we
 explored the link between taste sensitivity and taste liking. Only sour taste liking correlated with
- explored the link between taste sensitivity and taste liking. Onl
- ¹⁷ sucrose and by trend with sour thresholds.

18 Keywords: sensitivity; taste; threshold; staircase; QUEST; bias; quick Yes-No (qYN)

19 1. Introduction

The ability to taste is undoubtedly crucial for nutrient sensing. Yet, the ability to taste is scarcely assessed in large cohorts and precise and "practical" methods for the measurement of taste sensitivity are needed to get a better understanding of the extent to which taste shapes preferences and eating behavior.

The precise and reasonably quick measurement of sensory sensitivity has been a challenge in all senses. The relatively quick adaptation in gustation, however, increases the data collection burden enormously, as it requires long inter-stimulus intervals that cannot simply be countered with a reduction of the number of experimental trials. Using a Bayesian adaptive testing framework [3], we have previously shown that taste sensitivity can be precisely and reliably measured in a fraction of the time needed with conventional, non-adaptive methods [1,4].

Bayesian adaptive methods like QUEST [3] typically produce estimates of psychometric function parameters quicker than conventional staircase procedures, especially if the number of possible stimuli is large. This is mainly due to two features specific to Bayesian methods: firstly, they can incorporate prior knowledge by assigning a probability to each individual parameter value; and secondly, they

incorporate the entire response history of a participant to predict the "true" parameter value and select 34 the next stimulus. Accordingly, on every trial, stimuli are selected such that the expected knowledge 35 gain about the parameter (e.g., threshold) is maximized. This may lead to relatively strong intensity 36 changes from one trial to the next, especially at the beginning of an experimental run, and allows the 37 procedure to converge within fewer trials compared to a staircase. 38 When measuring thresholds, participants' responses not only depend on their sensory sensitivity, 39 but are also influenced by mental processes. Specifically, whether a participant will report the presence 40 of a stimulus is influenced by their cognitive *response criterion*: if a *liberal* criterion is adopted, even weak, non-obvious stimuli will be reported; if the criterion, however, is strict, only stimuli that the 42 participant believes are sufficiently strong are going to be reported. Generally, a liberal criterion leads 43 a higher false-alarm rate (FAR), which describes the proportion "yes" responses to a blank, while a 44 stricter criterion reduces the frequency of false alarms [5,6]. 45 Although QUEST was designed for alternative forced-choice tasks, which are commonly thought 46 to control the response criterion (but c.f. [7] for criticism of this view), the simplicity of the yes-no 47 experiment (see [7] for a systematic review of adaptive procedures) has been a strong motivator to 48 explore the suitability of QUEST in a yes-no design, and it has yielded good performance in the 49 chemosensory domain [1,4,8]. In these studies, participants were instructed to be "conservative" in 50 their response behavior, in an attempt to keep false-alarm rates low and constant across sessions. This 51 is imperative because QUEST can only estimate a single parameter, such that when the threshold is to 52 be estimated, all other parameters defining the psychometric function, like FAR, slope, and the lapse 53 rate (i.e. the proportion of "no" responses to high-intensity, supra-threshold stimuli) need to be set to a 54 fixed value *a priori*. While the lapse rate can safely be assumed to be low in taste threshold testing, 55 provided that sufficiently long inter-stimulus intervals are used and participants thoroughly rinse 56 their mouth between trials, FAR may vary between repeated measures despite our instructions. Slope determines how well participants can detect intensity differences between stimuli, and as such can also 58 serve as an (implicit) measure of threshold reliability. Just like thresholds differ between participants 59 and, obviously, tastants, it is known from other sensory systems that "slopes are different for different 60 stimuli, and this can lead to misleading results if slope is ignored." [7] 61

Here, we set out to address these concerns, and tested whether FAR and slope indeed vary between tastants and sessions, and whether the measurement of these parameters helps improve the repeatability of taste threshold measurements. To this end, we derived a test-retest design in which we employed two Bayesian procedures, the previously used QUEST, which only measures threshold, and qYN [9], which assesses the sensory threshold, FAR, and psychometric function slope.

67 2. Materials and Methods

68 2.1. Participants

41 participants (34 women; mean age M = 30.1, standard deviation SD = 11.4, range 18–64 years) participated in the study and received compensatory payment. Their weight was within the 70 normal range according to the body mass index (BMI; M = 22.5, SD = 2.6). Four participants were 71 smokers. Exclusion criteria were self-reported taste and smell disorders, current or recent oral, nasal or 72 sinus infections, pregnancy, recent (during the last 6 months) childbirth, metabolic disease, and recent 73 (during the last 3 months) weight change exceeding 10 kg. No participants were excluded, however, the samples for the different analyses vary slightly because individual data point were missing. The 75 age and sex structure for each sub-sample can be obtained in the associated data file (see Data and 76 software availability section). The study conformed to the revised Declaration of Helsinki and was 77 approved by the ethical board of the German Society of Psychology (DGPs). 78

79 2.2. Procedure

⁸⁰ 2.2.1. Experimental sessions

Participants were invited to four experimental sessions that lasted 1 hour each: a Test and a Retest session for each of two threshold algorithm. To ensure similar testing conditions across sessions, participants were instructed to refrain from eating and drinking anything but water 30 min before visiting the laboratory. Further, the sessions were scheduled at approximately the same time of day, and within 10 days (inter-session interval: M = 2.3, SD = 2.0, range 1–10 days).

At the beginning of the first session, participants completed a screening questionnaire, the Dutch Eating Behavior Questionnaire (DEBQ) [10], and rated how much they like and how often they typically consume salty, sour, sweet, and bitter foods. Following the ratings and in each subsequent session, taste recognition thresholds for citric acid (sour), sodium chloride (salty), quinine hydrochloride (bitter), and sucrose (sweet) were measured using either of two algorithms, QUEST or qYN, described below. The order of tastants was balanced across participants and kept constant for Test and Retest within each participant. The order of algorithms was balanced across participants.

⁹³ 2.2.2. Eating behavior, taste liking, and food consumption

Eating style was measured with the DEBQ [10], which assesses – along three dimensions – the

degree of restrained, emotional, and external eating behavior. Participants could chose between the

German and English version; 7 participants completed the English version. The questionnaire data of

one participant (female, 27 years old) are missing.

Furthermore, participants rated how much they liked salty, sour, sweet, and bitter foods andbeverages on separate, horizontal 5-point Likert scale anchored with 1 (not at all) and 5 (extremely).

They furthermore provided the frequency at which they typically consume salty, sour, sweet, and bitter foods and beverages on a scale with 7 options: daily (score 7), 4–6 times per week (6), 2–3 times per week (5), once per week (4), 2–3 times per month (3), once per month (2), less than once per month (1). Specifically, they provided ratings for the following items: sweet, sour, and bitter beverages, sweet, sour, and bitter fruits and vegetables, sweet cake/candy, salty snacks as well as added salt. Rating for each taste quality were averaged for further analysis. The ratings were assessed in paper–pencil format. Ratings from four participants (female, 21–27 years old) are missing.

¹⁰⁷ 2.2.3. Taste recognition thresholds

108 procedure

Participants were seated comfortably and blindfolded to reduce distraction and improve focus. At 109 the beginning of each measurement, they were told which taste would be tested next. At the beginning 110 of each trial they were asked to stick out the tongue and received the stimulus. Participants were 111 required to indicate whether they recognized the taste by nodding ("yes") or shaking their head ("no") 112 while they kept their tongue extended. Immediately after the response, the experimenter logged it 113 into the computer, and participants rinsed their mouth with deionized water. Participants received no 114 feedback as to their performance during the experiment. The interval between consecutive stimuli 115 was approx. 30 s. 116

117 Taste stimuli

Tastants were prepared by diluting prototypical chemicals that are known to elicit a clear taste perception in deionized water: citric acid (sour; molar mass $M = 192.12 \text{ g mol}^{-1}$), sodium chloride (salty; $M = 58.44 \text{ g mol}^{-1}$), quinine hydrochloride (bitter; $M = 396.91 \text{ g mol}^{-1}$), and sucrose (sweet; $M = 342.30 \text{ g mol}^{-1}$). All chemicals were produced by Sigma-Aldrich and purchased from Merck KGaA, Darmstadt, Germany. ¹²⁶ 0.273); quinine hydrochloride, 0.077×10^{-3} mM to 3.131 mM (21 log₁₀ steps; step width: 0.230); sucrose, ¹²⁷ 0.073 mM to 584.283 mM (14 log₁₀ steps; step width: 0.300).

Taste solutions were stored refrigerated at 4 °C for a maximum duration of seven days in glass bottles. During testing, they were sprayed manually by the experimenter to the anterior half of the tongue using a conventional spray head that released approx. 0.2 mL.

131 Psychometric functions

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The QUEST implementation we used assumes a psychometric function in which the proportion of "yes" responses to a stimulus is given by

$$\Psi_{\text{yes}}(c) = \delta \gamma + (1 - \delta) [1 - (1 - \gamma) \exp(-10^{\beta(c - \tau + \epsilon)})].$$

Here, c is the concentration of a given tastant, and τ is the threshold concentration one wishes 134 to estimate. The parameter ϵ shifts the psychometric function such that threshold performance 135 (proportion of "yes" responses) equals a predefined value; in our case, 0.8 or 80 %. ϵ was determined 136 automatically by our software based on the desired threshold performance. The parametrization of the 137 procedure was identical to the one used by [1]: We defined "threshold" as the concentration with an 138 expected proportion of 80% "yes" responses ($\Psi_{\text{ves}}(c) = 0.80$); the prior probabilities for the threshold 139 parameter were given by a normal distribution with a standard deviation of 20, centered on the starting 140 concentration of the respective tastant (see the Taste threshold stimulus selection section below). All other 141 parameters were fixed *a priori*: slope, β , to 3.5; and both the false-alarm and lapse rate, γ and δ , to 0.01. 142 Internally, QUEST works with an abstract "intensity grid", whose granularity we set to 0.01 as well. 143

In quick Yes-No [9], the psychometric function describing the proportion of "yes" responses for a given concentration is

$$\Psi_{\text{ves}}(c) = \epsilon + (1 - \epsilon) \left[1 - \Phi \left(\lambda - d'(c) \right) \right]$$

with Φ being the cumulative normal distribution with a mean of 0 and a standard deviation of 1. ϵ is the lapse rate, which describes how frequently stimuli of a high intensity are not recognized. The decision criterion, λ , is related to the FAR via the the percent point function of the normal distribution: $\lambda = \Phi^{-1}(1 - FAR)$ [9]. Ψ_{yes} depends on the sensitivity function, d', given by [9]

$$d'(c) = \frac{\beta(c/\tau)^{\gamma}}{\sqrt{(\beta^2 - 1) + (c/\tau)^{2\gamma}}}.$$

au is the "threshold intensity", which here is defined as the tastant concentration corresponding 150 to a sensitivity of d' = 1. β defines the upper asymptote and γ the slope. In this study, we 151 estimated threshold, τ ; slope, γ ; and decision criterion, λ . Consequently, the parameter space 152 was a three-dimensional a grid. Different ranges of τ were used for each tastant. To achieve a 153 finer granularity, additional (virtual) concentration steps were inserted halfway between the existing 154 (physical) concentration steps, producing 27 values for citric acid and sucrose, 23 for sodium chloride, 155 and 35 for quinine hydrochloride. For γ , we used 10 values in the interval [0.5, 3.0], evenly spaced 156 on a decadic logarithmic grid; and for λ , we used 8 evenly spaced values in the interval [0.75, 2.50], 157 corresponding to FARs of [22.7%, < 1.0%]. We assumed no prior knowledge regarding the "true" 158 parameter values, and, hence, used an "uninformative" prior that assigned the same probability to all 159 possible parameter value combinations. β was fixed at 5.0 [9], and ϵ was set to 0.

161 Stimulus selection

The concentration presented in the first trial for each tastant was predefined such that it would be 162 supra-threshold for most participants, in order to familiarize them with the particular tastant as testing 163 commenced (citric acid: 7.328 mM; sodium chloride: 97.469 mM; quinine hydrochloride: 0.077 mM; 164 sucrose: 73.509 mM). For subsequent trials, the QUEST and qYN procedures proposed the stimulus 165 concentration to present based on response behavior in all previous trials. While QUEST aims to place 166 stimuli at threshold concentration, qYN – trying to estimate *three* parameters at once – typically also 167 suggests to present stimulus concentrations slightly above and below threshold to measure slope, and 168 at very low concentrations to measure estimate the FAR. 169

As both algorithms internally worked with smaller, virtual concentration steps, they would 170 sometimes propose concentrations that were not physically available. In this case, our computer 171 program selected the concentration closest to the proposed one, and informed the algorithm about the 172 actually used concentration. In QUEST, we added an additional rule: whenever the algorithm proposed 173 to present the same concentration on two consecutive trials, we increased the concentration in the 174 second trial by one step if the participant had responded "no" to the previous trial, and we decreased the concentration by one step if the response in the previous trial was "yes", thus introducing a little 176 more variability to avoid repetitive presentation of the same concentration on multiple consecutive 177 trials, which we felt could have been more tiring for participants. In qYN, we did not add such a rule, 178 as the algorithm itself introduced somewhat abrupt concentration changes once in a while in order to 179 determine FAR and slope.

181 Taste recognition termination

qYN experimental runs always ended after 20 trials. For QUEST, we employed the same termination criterion as in a previous study [1]: after more than 10 trials had been performed, we checked after each trial whether the 90% confidence interval of the threshold estimate was smaller than half a concentration step; if that was the case, the experimental run was finished. Otherwise, a maximum of 20 trials were performed.

187 Parameter estimates

To retrieve the final threshold estimate, we calculated the mean of the concentrations weighted by the posterior distribution (QUEST) or marginal posterior distribution (qYN), respectively. This measure had been shown to produce an unbiased estimate of threshold in QUEST, as opposed to other metrics [11]. We limited the values of the threshold estimates to the range of stimulus concentrations used in the present study, as QUEST could – in rare cases – produce thresholds outside of this range for extremely sensitive or insensitive participants. The was the case for a single participant where QUEST produced one quinine hydrochloride threshold above the highest available concentration.

For qYN, FAR, and slope were calculated as the mean of the respective parameter space weighted by the corresponding marginal posterior.

197 2.3. Analysis

The significance level α was set to 0.05 *a priori* for all statistical tests. Greenhouse-Geisser correction was applied for violation of sphericity in repeated measures analysis of variance (rmANOVA); uncorrected degrees of freedom and corrected *p*-values are reported in this case.

201 2.3.1. Ratings

Ratings for taste liking and frequency of consumption were submitted to separate one-factorial
rmANOVA with four levels (sour, salty, sweet, bitter). To quantify the potential link between taste
liking and sensitivity, we computed Spearman's correlation coefficients. For this, the average of all
threshold estimates from QUEST and qYN for a given participant was used as robust measure for

taste sensitivity. Data from only 37 participants are reported for liking and frequency of consumption
 because 4 participants did not complete the ratings.

208 2.3.2. Taste recognition data cleaning

Out of the 656 obtained thresholds (41 participants × 4 tastants × 2 procedures × 2 sessions), 41 (6.3%; 17 QUEST and 24 qYN) were lost during the transfer of electronic data, yielding 615 datasets. Because in many instances only data from a single session was lost, we used the measurements for which data from both sessions was available. The resulting 590 datasets entered analysis (90% of the total data; 302 QUEST and 288 qYN runs). All included and omitted datasets are available online (see the *Data and software availability* section for details).

- 215 2.3.3. Test-retest reliability
- 216 Threshold

²¹⁷ We first submitted the threshold estimates to separate rmANOVAs for each tastant with the ²¹⁸ factors *procedure* (QUEST, qYN) and *session* (Test, Retest) to test for systematic differences between ²¹⁹ procedures and measurement repetitions.

We then calculated Spearman's rank correlation, ρ , for each tastant to quantify the monotonic relationship between the measurement results in both sessions.

Correlation analysis does not necessarily provide a good indication of absolute *repeatability* of an 222 experiment, as "[the] correlation coefficient is a reflection of how closely a set of paired observations 223 (test-retest data in this case) follow a straight line, regardless of the slope of the line" [12], and it 224 also disregards systematic changes between measurements [12,13], such as a constant offset. We, 225 therefore, conducted an additional analysis that focuses on the *differences* between measurements. For 226 each procedure separately, we first calculated the difference between Test and Retest estimates for 227 all participants; then, we calculated the standard deviation of these differences, s_d , and derived a 228 coefficient of repeatability (CR), CR = $1.96 \times s_d$ [14]. ¹ The number 1.96 is a *z*-score and corresponds 229 to the 97.5% quantile of the normal distribution. If a participant were measured repeatedly using the 230 same procedure, we would then expect 95 % of the absolute measurement differences not to exceed 231 CR. This provides a straightforward, single-number representation of the magnitude of measurement 232 variation to expect. Lastly, we calculated the mean of the differences between sessions, d, and estimated 233 the 95 % limits of agreement (LoA) as LoA = $d \pm CR$ [14]. These limits correspond to the 95 % confidence interval of the differences, and, consequently, narrower LoAs suggest better measurement repeatability. 235 Because the calculations of the mean difference and LoAs are based on an experimental sample, 236 they are estimates that naturally have a certain amount of uncertainty associated with them. We 237 therefore also derived 95% confidence intervals (CIs) of these estimates. The mean difference was 238 assumed to be normally distributed with mean \bar{d} and SD s_d/\sqrt{n} ([2]; with the number of paired samples, *n*), and hence the CI corresponded to the 2.5 % and 97.5 % quantiles of this distribution. CIs 240 of the LoAs were calculated via the "exact paired" method [15]. 241

For visual comparison, we plotted the differences between Test and Retest over the mean of both measurements (which serves as our best estimate of the "true" value) and added the mean difference \bar{d} and LoAs as horizontal lines, producing so-called *Bland-Altman* or *Tukey mean difference plots*. These plots allow for a quick and straightforward inspection of measurement differences, exposing systematic biases ($\bar{d} \neq 0$) the degree of measurement differences and their variability.

¹ Note that it has been suggested [2,12,13] to calculate CR as $\sqrt{2} \times 1.96 s_w$, where s_w is the *within-participant standard deviation*, i.e., the square-root of the averaged within-participant variances of measurement repetitions. We found that with our data, this approach produced very similar results (deviating only in the second decimal place) to the simpler formula $1.96 \times s_d$, which directly and intuitively corresponds to the 95 % limits of agreement in the Bland-Altman plots. Therefore, we elected to follow this simpler approach, and report CRs based on the SD of measurement differences between sessions here.

247 False-alarm rate (FAR)

The decision criterion parameter λ , which was only estimated by qYN, was first transformed to a proportion of false alarms by inverting the relationship given in section 2.2.3.3: FAR = $1 - \Phi(\lambda)$. The FARs were submitted to an rmANOVA with the factors *tastant* (citric acid, sodium chloride, quinine hydrochloride, sucrose) and *session* (Test, Retest) investigate whether different tastants and measurement repetitions affected FARs differentially.

We then constructed Bland-Altmann plots as described above. The plots revealed that the the 253 difference between both sessions changed with the magnitude of the session mean. However, the 254 calculation of CRs and LoAs requires the differences between sessions to be approximately normally 255 distributed to work correctly. If that is not the case, a logarithmic transformation can be carried 256 out prior to analysis [13]. Here, we elected to use the decadic logarithm, \log_{10} . After the data had 257 been transformed, d, CR, LoAs, and CIs were calculated following the same procedure as described 258 above for thresholds. To plot the results that were calculated in log space, x_{log} , in their original 259 coordinate system for intuitive visual assessment, a back-transformation must be applied [16]: $y_{\text{back}} =$ 260 $2M(10^{y_{log}}-1)/(10^{y_{log}}+1)$, where M is a given value on the abscissa (i.e., any given session mean). 261 The result of the back-transformation, therefore, describes a line, not a single value, in the original 262 coordinate system. In other words: the value of the back-transformed parameter, x_{back}, is conditional 263 on M. The generated plots can be interpreted in the same way as a Bland-Altman plot that did not 264 undergo a logarithmic transformation. 265

266 Slope

Similarly to FARs, the slopes estimated by qYN were analyzed using rmANOVA with the factors
 tastant and *session*, and Bland-Altmann plots, CRs, LoAs and their CIs were derived as described above
 for the threshold data.

270 Relationship between FAR and slope

As the FAR determines the location of the lower asymptote, changes in FAR necessarily lead to changes in steepness of the psychometric function if threshold is assumed to remain unchanged: as FAR increases, the slope must decrease, and vice versa. We thus calculated Spearman's rank correlation between FARs and slopes, pooled across all tastants, in order to explain potential fluctuations of FAR and slope across sessions, which could be explained through the covariance of both variables.

276 2.3.4. Software

Stimulus presentation and data collection were guided by a Python computer program based on
PsychoPy 1.85.4 [17] on Windows 7 (Microsoft Corp., Redmond, WA/USA). Statistical analyses were
carried out with JASP 0.11.1 (https://jasp-stats.org/) and pingouin 0.2.9 [18]. CIs for the Bland-Altman
plots were calculated via pyCompare (https://github.com/jaketmp/pyCompare). Plots were created
using matplotlib 3.1.1 (https://matplotlib.org) and seaborn 0.9.0 (https://seaborn.pydata.org).

282 3. Results

Measure	Mean	SD	Ν
DEBQ emotional eating	2.22	0.64	40
DEBQ restrained eating	2.55	0.67	40
DEBQ external eating	3.02	0.65	40
Liking of salty	3.41	1.05	37
Liking of sour	3.05	1.09	37
Liking of sweet	4.11	0.83	37
Liking of bitter	1.89	0.98	37
Consumption of salty	3.64	1.33	37
Consumption of sour	4.19	1.10	37
Consumption of sweet	4.70	1.03	37
Consumption of bitter	4.00	1.24	37

Table 1. Questionnaire and rating data.

DEBQ and liking scores ranged from 1 to 5. Consumption frequencies ranged from 1 to 7.

283 3.1. Eating behavior, taste liking, and food consumption

All measurement outcomes are summarized in Tab. 1.

Participants exhibited eating style scores that conform well with recent norm data [19] and can,
hence, be considered normal in eating behavior. Note that the questionnaire data of one participant
(female, 27 years old) was lost.

The liking of different tastes varied significantly ($F_{3,36} = 33.68$, p < 0.001, $\eta_p^2 = 0.483$), as expected, 288 with bitter taste being significantly less liked than salty ($t_{36} = -6.95$, $p_{holm} < 0.001$), sour ($t_{36} = -7.17$, 289 $p_{\text{holm}} < 0.001$), and sweet ($t_{36} = -9.65$, $p_{\text{holm}} < 0.001$); and with sweet being significantly more liked 290 than salty ($t_{36} = 2.77$, $p_{holm} = 0.018$) and sour ($t_{36} = 4.2$, $p_{holm} < 0.001$). Accordingly, scores (min=1, 291 max=5) were highest for sweet, followed by salty, sour, and bitter. In contrast to sweet, sour, and 292 salty, which were generally liked (as indicated by ratings larger than 3), bitter was clearly disliked (as 293 indicated by a rating smaller than 3). The liking of sour taste correlated significantly with the sucrose 294 ($\rho = 0.403$, p = 0.013) and marginally with the citric acid ($\rho = 0.305$, p = 0.066) taste threshold. No 295 further correlations between taste liking and thresholds were found (all p > 0.19). 296

Participants reported to consume food and beverages of the different tastes only approximately
 once per week (score=4) for all taste qualities). Because the reported mean frequencies raise doubts as
 to their validity, no further statistical analyses were conducted.

300 3.2. Taste recognition

The distributions of threshold estimates, averaged across both sessions and split by procedure, 301 are shown in Fig. 2. The rmANOVAs revealed a main effect of *procedure* – indicating that threshold 302 estimates were systematically lower for qYN compared to QUEST – for citric acid ($F_{1,32} = 17.475$, 303 $p < 0.001, \eta_p^2 = 0.353$), sodium chloride ($F_{1,32} = 44.728, p < 0.001, \eta_p^2 = 0.583$), and sucrose 304 $(F_{1,31} = 11.198, p < 0.01, \eta_p^2 = 0.265)$, but not for quinine hydrochloride $(F_{1,33} = 3.241, p = 0.08, p < 0.01)$ 305 $\eta_p^2 = 0.089$). Thresholds did not differ significantly between sessions for sodium chloride ($F_{1,32} = 0.342$, 306 p = 0.56), quinine hydrochloride ($F_{1,33} = 1.195$, p = 0.28), and sucrose ($F_{1,31} = 0.219$, p = 0.64); 307 however, we found a main effect of session for citric acid ($F_{1,32} = 4.492, p = 0.042, \eta_p^2 = 0.123$). No 308 interactions between *procedure* and *session* were found (all p > 0.13). The threshold estimates, their 309 respective minimum and maximum values, and their standard deviations are listed in Tab. 2. Graphical 310 representations of the psychometric functions, generated based on the mean parameter estimates (i.e., 311 averaged across participants), are shown in Fig. 1. Figures of individual psychometric functions are 312 available online; see the Data and software availability section for details. 313

On average, QUEST needed 14.6 trials – corresponding to 6:50 min – to converge to a threshold. For qYN, the duration of an experimental run was always approx. 9:30 min, as the number of trials was fixed to 20.

				Threshold in log ₁₀ mM			
procedure	Tastant	Ν	Session	mean	min	max	SD
QUEST	Citric Acid	37	Test	-0.141	-1.564	1.350	0.621
			Retest	-0.256	-1.540	1.385	0.666
	Sodium Chloride	38	Test	1.140	-0.417	2.495	0.631
			Retest	1.069	-0.432	2.100	0.544
	Quinine-HCl	38	Test	-1.737	-3.514	0.496	1.101
			Retest	-1.889	-3.514	0.339	0.953
	Sucrose	38	Test	1.054	-0.592	2.414	0.705
			Retest	1.089	-0.563	2.194	0.660
qYN	Citric Acid	37	Test	-0.446	-1.807	0.812	0.563
			Retest	-0.558	-1.508	0.574	0.598
	Sodium Chloride	36	Test	0.785	-0.457	2.052	0.607
			Retest	0.831	-0.369	2.222	0.571
	Quinine-HCl	35	Test	-1.974	-3.409	0.239	0.952
			Retest	-1.980	-3.369	0.091	0.962
	Sucrose	36	Test	0.871	-0.613	2.243	0.672
			Retest	0.765	-0.742	1.881	0.669

Table 2. Results of the threshold measurements during Test and Retest for QUEST and qYN.

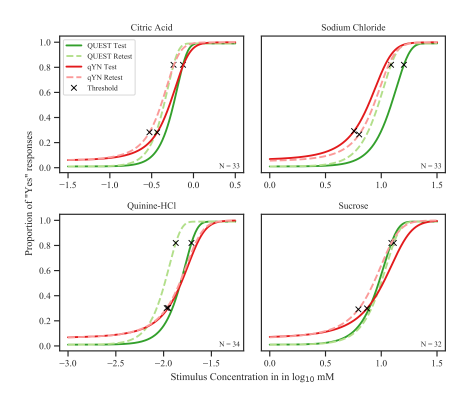


Figure 1. Psychometric functions based on the averaged parameter estimates. Mean thresholds are depicted as crosses. It is obvious how the definition of "threshold" differs between QUEST and qYN: while threshold in QUEST is solely based on the proportion of "yes" responses, qYN uses a threshold based on the sensitivity function d', which also takes false alarms into account. Only data from participants for whom threshold data of both sessions and procedures was available is included to facilitate visual comparison.

317 3.2.1. Threshold repeatability

Test and Retest threshold estimates correlated significantly for all tastants in both QUEST (citric 318 acid: $\rho_{35} = 0.62$, sodium chloride: $\rho_{36} = 0.63$, quinine hydrochloride: $\rho_{36} = 0.80$, sucrose: $\rho_{36} = 0.67$; 319 all p < 0.01) and qYN (citric acid: $\rho_{35} = 0.71$, sodium chloride: $\rho_{34} = 0.60$, quinine hydrochloride: 320 $\rho_{34} = 0.76$, sucrose: $\rho_{33} = 0.76$; all p < 0.01); see Fig. 3. To gain a better understanding of the nature of 321 the individual differences between Test and Retest thresholds, we constructed Bland-Altman plots for 322 each tastant in both procedures (Fig. 4). The 95% confidence intervals of the mean differences always 323 included 0, providing no evidence of systematic differences between sessions. We then derived the 324 coefficients of repeatability (CR); the results of two measurements would be expected to differ no more 325 than CR in 95% of the cases. The respective CRs for QUEST and qYN were, in log₁₀ mM: 0.97 and 0.84 326 for citric acid (corresponding to 3.6 and 3.1 concentration steps), 0.98 and 0.95 for sodium chloride (3.6 327 and 3.5 steps), 1.07 and 1.29 for quinine hydrochloride (4.7 and 5.6 steps), and 1.10 and 1.03 for sucrose 328 (3.7 and 3.4 steps). Accordingly, the mean CR across all tastants was 3.90 for both procedures. 329

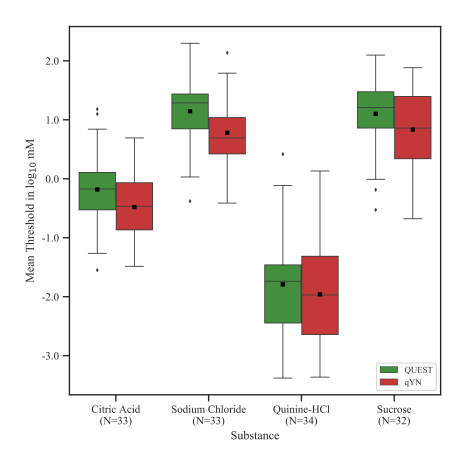
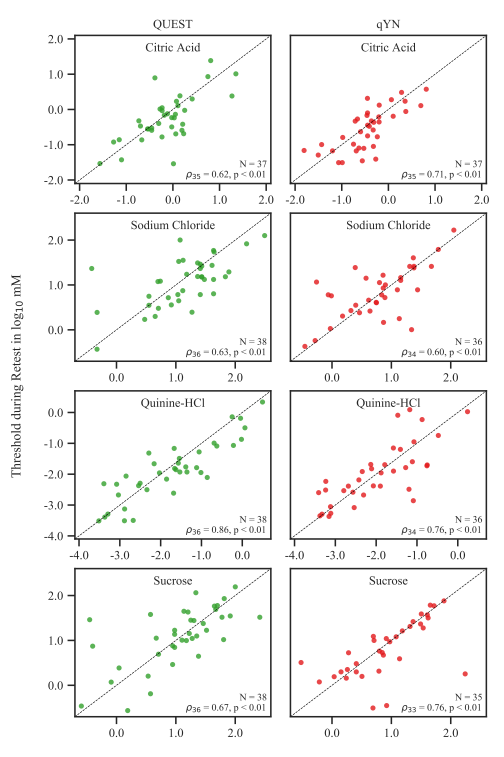
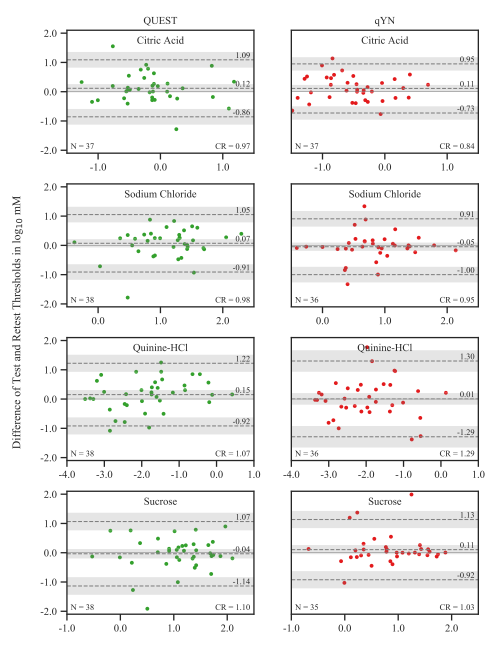


Figure 2. Distributions of the means of Test and Retest threshold estimates, split by tastant and procedure. Squares indicate the mean, and whiskers correspond to $1.5 \times$ inter-quartile range. Only participants for whom threshold data for both sessions and procedures was available are included; the number of participants is given below the respective abscissa labels for each tastant.



Threshold during Test in log10 mM

Figure 3. Correlation between Test and Retest threshold estimates for QUEST and qYN. Each point represents one participant; the dashed line is the identity line.



Mean of Test and Retest Thresholds in log10 mM

Figure 4. Bland-Altman plots showing differences between Test and Retest thresholds for QUEST and qYN, mean difference \bar{d} , and limits of agreement (LoA) corresponding to 95% confidence intervals (CIs) as $\bar{d} \pm 1.96 \times$ SD. The shaded areas represent 95% CIs of these estimates.

330

To investigate potential differences between tastants and across sessions in estimated FARs and psychometric function slopes, we conducted two rmANOVAs with the factors *tastant* and *session*, with FAR and slope as the respective dependent variables.

For FARs, we found no main effects and no interaction of the factors (all p > 0.25), indicating 334 there was no evidence that response criteria would systematically vary with tastants or across sessions. 335 We therefore decided to pool all data points, yielding a mean FAR of 0.059 (SD=0.024) spanning across 336 a relatively wide range (0.017–0.216). Yet, only a single of the 37 participants that had entered analysis 337 showed FARs > 0.10 in *all* measurements. The Bland-Altman plot revealed an increasing variability of 338 the differences between Test and Retest as the magnitude of FARs increased (Fig. 5). This finding can be 339 interpreted such that some participants would expose a relatively high FAR in one session, but a small 340 FAR in the other. For the majority of participants, however, FARs varied within a relatively narrow 341 range. Because the FAR differences between sessions were obviously not normally distributed, we 342 log₁₀-transformed the data for the calculations of mean difference, CR, LoAs, and the corresponding 343 CIs. The results were then back-transformed to the original scale of the data [16]. As can be seen in Fig. 5, the back-transformation does not yield a single value, but a line spanning across the session 345 means. The resulting CR was $0.969 \times \text{session mean}$. 346

For d' slopes, we found a significant main effect of *tastant*, albeit with a small effect size ($F_{3,96}$ = 347 3.05, p = 0.04, $\eta_p^2 = 0.09$). This finding suggests that the ability to discriminate between stimuli 348 of adjacent concentration steps systematically shifted with the presented tastants. A post-hoc *t*-test revealed that this effect was driven by a significant difference between sodium chloride and sucrose 350 slopes ($t_{33} = 2.857$, $p_{holm} = 0.045$, d = 0.497). There was no effect of *session* and no interaction between 351 the factors (both p > 0.19). Mean slopes pooled across sessions were 1.49 (SD=0.53) for citric acid, 1.69 352 (SD=0.60) for sodium chloride, 1.46 (SD=0.57) for quinine hydrochloride, and 1.45 (SD=0.46) for sucrose. 353 Bland-Altman plots for all tastants are shown in Fig. 6. In agreement with the rmANOVA results, differences between sessions did not significantly deviate from zero, as indicated by the confidence 355 intervals spanning across 0. CRs ranged from 0.99 (sucrose) to 1.32 (sodium chloride), which is large, 356 considering the mean slopes. 357

There was a significant correlation between FARs and slopes ($\rho_{286} = -0.749$, p < 0.001), indicating that higher FARs were associated with reduced steepness of the d' sensitivity function.

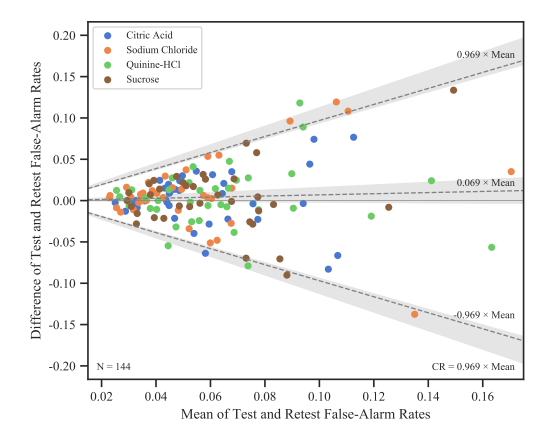
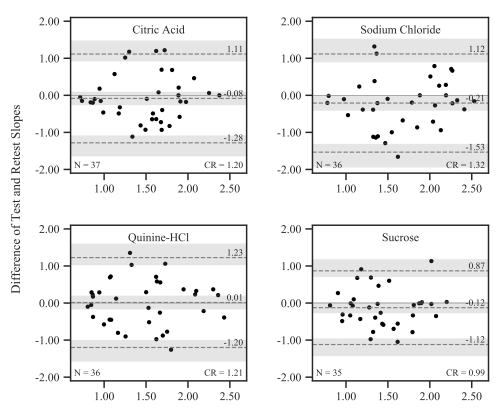


Figure 5. Bland-Altman plot for qYN false-alarm rates (FAR). Markers show differences between Test and Retest FARs for all available Test-Retest pairs from all 37 participants . Since the variability of session differences increases with session means, the data was

Since the variability of session differences increases with session means, the data was \log_{10} -transformed before calculating mean difference \bar{d} , limits of agreement (LoA) corresponding to 95% confidence intervals (CIs) as $\bar{d} \pm 1.96 \times$ SD, and 95% CIs of these estimates (shaded areas). The figure shows the *back-transformed* parameter values, plotted on the original scale of the data. Due to the back-transformation, the lines representing \bar{d} and LoAs have a slope $\neq 0$, i.e., they are not parallel to the abscissa, and we provide their respective formulas. Note that the intercepts of all lines were 0, and are therefore omitted.



Mean of Test and Retest Slopes

Figure 6. Bland-Altman plots showing differences between the slopes of the estimated sensitivity functions (*d'*) in qYN Test and Retest, mean difference \bar{d} , and limits of agreement (LoA) corresponding to 95% confidence intervals (CIs) as $\bar{d} \pm 1.96 \times$ SD. The shaded areas represent 95% CIs of these estimates.

360 4. Discussion

Using two Bayesian procedure, based on QUEST [3] and qYN [9], we explored the impact of false-alarm rate and psychometric function slope estimation on the precision and accuracy of taste sensitivity measurements.

4.1. Taste recognition thresholds

The comparison of the session means for each procedures showed slightly, but systematically higher thresholds for QUEST compared to qYN for citric acid, sodium chloride, and sucrose; the difference was not significant for quinine hydrochloride, which we believe can be attributed to the larger variability of bitter threshold measurements compared to the other taste qualities. Since the exact shape and parametrization of the psychometric function and the definition of "threshold" differed between both procedures, we expected the estimated threshold values to differ.

371 4.2. Threshold repeatability

Both, QUEST and qYN thresholds showed good monotonic relationships across sessions, as 372 indicated by test-retest correlations ranging from $\rho = 0.62$ to $\rho = 0.86$ for QUEST and from $\rho = 0.60$ 373 to $\rho = 0.76$ for qYN. No procedure produced consistently higher correlations than the other: qYN 374 correlations were stronger for citric acid and sucrose, while QUEST showed higher correlations for 375 sodium chloride and quinine hydrochloride. These values compare very well with a past applications 376 of the QUEST method (r = 0.59-0.83 [1]), with a modified Harris-Kalmus (r = 0.70-0.77 [20]), and 377 a forced-choice staircase procedure (r = 0.76-0.86 [21]). The latter two approaches required notably 378 longer testing times than QUEST. Other researchers have also observed much smaller correlations, 379 for example with the relatively quick three-drop method (r = 0.36-0.61 [22]) and with taste strips 380 (r = 0.34 - 0.56 [22]).381

The observed correlation coefficients do not, however, account for systematic changes occurring 382 between sessions, and do not necessarily honor the spread of the data and the slope associated 383 with their relationship [12]. Therefore, we a) created Bland-Altman plots [2,13,23] to visualize the 384 distributions of differences; and b) calculated coefficients of repeatability (CR; [14]) as an estimate of the expected measurement differences between sessions in an individual participant. We found 386 that thresholds did not vary systematically across sessions. qYN produced smaller CRs than QUEST 387 indicating better agreement between measurement repetitions – for all tastants except quinine 388 hydrochloride. The CR averaged across tastants was identical for both procedures at an equivalent of 389 3.90 concentration steps. Using a QUEST procedure in a 3-AFC task to estimate smell thresholds [8], we previously observed a CR corresponding to approx. 5.3 concentration steps on a \log_{10} grid with a 391 step width of 0.300, which is similar to the step width used here for sucrose, and larger than the step 392 width for all other tastants. Neglecting the task differences (yes-no in the present study versus 3-AFC 393 in [8]), the QUEST procedure seems to perform better for taste than for smell measurements. 394

We would like to emphasize a discrepancy between the estimated correlation coefficients and CRs. In QUEST, the highest correlation was found for quinine hydrochloride; yet, repeatability according to CR was better for sodium chloride and citric acid. In fact, repeatability was *best* for citric acid, yet the corresponding correlation was the *worst* of all of the four tastants. Similarly, in qYN the highest correlations were found for quinine hydrochloride and sucrose, but the respective CRs were highest, i.e. repeatability was lowest, for these tastants. The highest repeatability in the entire study was found for citric acid in qYN, yet the associated correlation was only found to be in a medium range.

Correlation coefficients are commonly adopted to quantify repeatability in the chemical senses
literature, and should therefore be calculated to enable comparisons with previously published studies.
Yet, thorough examination of the correlated data is required to ensure that the conclusions drawn
from these analyses are not inadvertently erroneous. We, therefore, suggest to always visualize the
data in a scatter plot and the identity line to uncover systematic changes between measurements,

which can occur even if the data points are highly correlated. In order to better understand the spread
and pattern of measurement differences, Bland-Altman plots and coefficients of repeatability (CR)
should be derived [2,12–14,23]. CRs indicate the magnitude of differences to expect when applying an
experimental procedure repeatedly, and help guide the decision whether that procedure is suitable for
a particular investigation, e.g., a clinical assessment.

412 4.3. False-alarm rates and d' slopes

False-alarm rates (FARs) and *d'* slopes were assessed in qYN runs. FARs did not differ between tastants, indicating that participants' response criterion was not taste-specific. Furthermore, FARs were generally low, suggesting that most participants complied with the instruction to be conservative in their response behavior. This finding was further substantiated through inspection of FAR differences between Test and Retest, which revealed little variation in most participants. Yet, for a few participants, the variance of session differences grew as the magnitude of FARs increased, leading to a relatively high FAR in one session, but a much smaller FAR in the other.

Slopes of the sensitivity function (*d'*) only differed between sodium chloride and sucrose. CRs for slopes were relatively large. While this parameter of the psychometric functions is known to be notoriously difficult to estimate, especially if the number of trials is small [9], slope and FAR are also directly linked: for a given sensitivity threshold, a lower FAR will lead to a steeper psychometric function while a higher FAR demands a reduction in steepness. We found evidence for this dependence, as FAR and slope were strongly negatively correlated.

Differences in FAR between sessions can be interpreted such that participants followed different 426 cognitive strategies in the two measurements, i.e., they changed their response criterion. These changes 427 in FAR, then, would inevitably affect the slope as well and *vice versa*. Inspection of the trial sequences 428 of experimental runs with the highest FARs (in the 90th percentile and above) revealed that, here, 429 participants had indeed responded "yes" to stimuli of very low concentrations that were clearly below 430 threshold. This shows that it is more likely that FAR changes lead to slope adjustments, than vice versa. 431 Overall, the results support our premise in the QUEST procedure that FARs are low and stable in the 432 majority of participants. 433

434 4.4. Measurement duration

On average, QUEST finished 2:40 min quicker than qYN, thanks to its dynamic termination criterion that ends the experimental run when the confidence interval around the threshold estimate reaches a predefined low limit. qYN, on the other hand, always completes 20 trials because no termination criterion was set in order to ensure sufficient data for the simultaneous estimation of the three parameters, threshold, FAR, and slope. Whether the amount of testing time required for qYN could be reduced by employing a similar dynamic stopping criterion needs to be tested in future studies.

442 4.5. Taste sensitivity, taste liking, and food preference

Additionally to taste thresholds, we assessed taste liking. The data revealed no clear link between 443 taste liking and taste sensitivity with the exception of a positive association between sour liking and 444 sucrose and by trend sour threshold. Accordingly, sour liking was higher in participants with lower sucrose and citric acid sensitivity (higher threshold). Whether this association has the potential to 446 shape taste and food preferences and food intake remains unanswered in the present study, as the 447 reported food frequencies appeared to be unrealistically low and could, therefore, not be used for 448 further analysis. A recent study, however, in which food and beverage consumption was thoroughly 449 assessed [24], showed clear associations between salty, sweet, and bitter taste sensitivity and the intake 450 of foods and beverages with these taste qualities. In this study, higher sensitivity was linked with lower 451 food intake (or vice versa). No such link was found for sour taste, though, leaving it open whether 452 taste sensitivity should be considered a general predictor for food intake behavior. The observation 453

that a low sour taste liking and poor sour taste abilities improved through the course of a weight loss
intervention obese children, indicates, however, that dietary changes may influence preference as well
as taste function, at least to some extent [25]. The latter data provide a glimpse into the complex and
potentially reciprocal interplay of taste function, taste preference, and food intake and a review on
the determinants of fruit and vegetable consumption revealed that other factors such as age, gender,
socio-economic status, preferences, parental intake, and availability play a major role as well [26],
thereby corroborating the multifaceted nature of food intake behavior and highlighting the need for
further studies.

462 5. Conclusions

We compared the repeatability of taste recognition threshold estimates produced by two adaptive 463 procedures, QUEST and qYN, for citric acid, sodium chloride, quinine hydrochloride, and sucrose. Both procedures select stimulus concentrations such that - based on a participant's entire response 465 history – the expected information gain about the true parameter(s) of a psychometric function is 466 maximized. While QUEST only assesses the threshold, qYN also adjusts FAR and slope. Our analysis 467 consisted of the widely adopted calculation of correlation coefficients between repeated measurements, 468 and the estimation of coefficients of repeatability (CR) to assess the expected difference between two measurements of the same participant. The magnitudes of test-retest correlations were generally good 470 and not clearly in favor of either threshold procedure. The CRs, however, revealed slightly better 471 repeatability of qYN thresholds for citric acid, sodium chloride, and sucrose, compared to QUEST. The 472 good agreement between both methods together with the low FARs observed in qYN suggest that, 473 overall, participants applied a conservative response criterion as instructed. 474

6. Data and software availability

The data reported in this paper along with graphical representations of individual threshold runs and psychometric functions are available for download from https://doi.org/10.5281/zenodo.3540534.

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