This is the author’s final peer reviewed version of the item published as:


Copyright : 2007, The Author
A complex relationship among chemical concentration, detection threshold, and suprathreshold intensity of bitter compounds

Russell SJ Keast (PhD) and Jessica Roper

The corresponding author may be contacted at the following address:

Dr. RSJ Keast
Senior Lecturer
School of Exercise and Nutrition Sciences
Deakin University
221 Burwood Highway, Burwood, Victoria 3125 Australia.
Phone: 03 9244 6944 International: +61 3 9244 6944
Fax: 03 9244 6017 International: +61 3 9244 6017
e-mail: russell.keast@deakin.edu.au
Abstract

Detection thresholds and psychophysical curves were established for caffeine, quinine-HCl (QHCl), and propylthiouracil (PROP) in a sample of 33 subjects (28 female mean age 24 ± 4). The mean detection threshold (±S.E.) for caffeine, QHCl and PROP was 1.2 ± 0.12 mM, 0.0083 ± 0.001 mM, and 0.088 ± 0.07 mM respectively. Pearson product-moment analysis revealed no significant correlations between detection thresholds of the compounds. Psychophysical curves were constructed for each bitter compound over six concentrations. There were significant correlations between incremental points of the individual psychophysical curves for QHCl and PROP. Regarding caffeine, there was a specific concentration (6mM) below and above which the incremental steps in bitterness were correlated. Between compounds, analysis of psychophysical curves revealed no correlations with PROP, but there were significant correlations between the bitterness of caffeine and QHCl at higher concentrations on the psychophysical curve (p<0.05). Correlation analysis of detection threshold and suprathreshold intensity within a compound revealed a significant correlation between PROP threshold and suprathreshold intensity (r=0.46-0.4, p<0.05), a significant negative correlation for QHCl (r=-0.33 to -0.4, p<0.05), and no correlation for caffeine. The results suggest a complex relationship between chemical concentration, detection threshold, and suprathreshold intensity.

Keywords

Bitter taste, caffeine, propylthiouracil (PROP), individual differences, threshold
Introduction

Taste receptors located on taste cells in the surface regions of our oral cavity are activated when chemicals enter our mouths. An electrical impulse is initiated and transferred via afferent fibers to cortical levels of the brain where it is decoded and we experience a perception associated with the chemical. A taste quality is experienced when the chemical concentration in the oral cavity reaches a level that not only activates a receptor, but the signal sent from the receptor is strong enough to elicit a perception. For example, a chemical may be in solution at a concentration that the sample population could not detect. As the concentration of the chemical increases a detection threshold will be reached, the level at which the chemical in solution may be discriminated from water. As the concentration of the chemical increases further the recognition threshold is reached, the point at which the quality (e.g., bitter) can be identified. As the concentration of the chemical increases still further, the intensity of bitterness mutually increases to a theoretical asymptote where concentration increases no longer cause subsequent increases in intensity (Keast & Breslin, 2003) (Figure 1).

Intuitively you may expect an individual with low detection threshold (sensitive to the chemical) to experience higher intensities at higher concentrations of the chemical compared to a second individual with higher detection threshold (insensitive to the chemical). An example of this intuitive model is observed with phenylthiocarbamide (PTC), if you have a low detection threshold for PTC (sensitive) you will be sensitive throughout the entire psychophysical function for that compound (Bufo et al., 2005). However, such relationships are not the norm (Bartoshuk, 2000; Mojet et al., 2003) presumably due to both genetic and environmental factors influencing bitter taste, and the complex nature of the organization of the oral peripheral and central cognitive system involved in bitter taste processing.
There is a large family of approximately 30 putative bitter-taste receptors (TAS2R’s) (Adler et al., 2000; Chandrashekar et al., 2000) located on bitter taste cells (Mueller et al., 2005). There are also many post-receptor transduction mechanisms including α-gustducin (McLaughlin et al., 1992), a phospholipase β subtype (Rossler et al., 1998), and transient receptor potential ion channels (Perez et al., 2002) to name a few. Any one bitter compound may access multiple transduction mechanisms. For example, caffeine is capable of translocating through cellular membranes and accessing 2nd messenger systems associated with bitter taste (Peri et al., 2000) and quinine-HCl (QHCl) can also activate non-receptor mechanisms associated with bitter taste cells (Caicedo et al., 2003; Kinnamon & Cummings, 1992; Rosenzweig et al., 1999). While there are multiple mechanisms on or within the bitter taste cell, the bitter quality we perceive is controlled by the taste cell, not the receptors; TAS2R’s expressed on sweet taste cells confer appetitive quality to what should be an aversive chemicals (Mueller et al., 2005).

An electrical signal leaves the taste cell and is transferred via afferent fibers to the subcortical areas nucleus of the solitary tract, followed by the second order synapse in the thalamus, before terminating in several regions of the insula (important in detection and suprathreshold intensity), frontal operculum cortex, and the orbital frontal cortex (important in hedonics). As the signal progresses upstream towards the cortical regions of the brain, greater selectivity of activation is observed and the neurons in the orbital frontal cortex may respond to only one taste quality. The cortical and sub-cortical regions of the brain integrate the signals and introduce plasticity into the gustatory system with feed-forward and feed-back pathways in operation (Jones et al., 2006; Katz et al., 2002).
Complexity of bitter taste

Presumably, differences in the quality and quantity of the multiple cellular mechanisms associated with bitter taste cells manifest in the large individual variation observed in bitter taste perception (Bartoshuk et al., 1998; Delwiche et al., 2001; Keast & Breslin, 2002b; Yokomukai et al., 1993).

Even though there is large variation in bitter taste perception, there is some commonality to bitter taste elicited by multiple chemicals, and these associations have been supported in human psychophysical studies (Delwiche et al., 2001; Keast & Breslin, 2002a; Lawless, 1979; McBurney, 1969). The most studied of all bitter chemicals that have commonality of bitterness are propylthiouracil (PROP) and PTC, primarily because there is known heritable variability in bitter taste perception which is related to halotypes of the TAS2R38 gene (Duffy et al., 2004; Bufe et al., 2005). Other bitter compounds such as caffeine and QHCl have also been extensively studied and commonality in suprathreshold bitterness has been established by phenotypic variation and genetic modeling (Hansen et al., 2006). However, there is no commonality between PROP bitterness and the bitterness elicited by QHCl and caffeine (Delwiche et al., 2001; Hansen et al., 2006; Keast et al., 2003a).

In the present study, the objective was to assess the relationship between chemical concentration, detection threshold, and suprathreshold intensity within and between three bitter compounds. Caffeine and QHCl were selected as they share commonality in suprathreshold bitterness perception and therefore may have commonality at detection thresholds level. PROP was selected as it elicits bitterness independent of caffeine and quinine, and the bitterness of PROP has been linked to a single receptor, TAS2R38.
Materials and Methods

SUBJECTS

Subjects (n=33, 23±4 years old, 28 female) between the ages of 18 and 38 were University students in Melbourne, Australia. All subjects agreed to participate and provided informed consent on an approved Institutional Review Board form. The participants, all non-smokers, were asked to refrain from eating, drinking or chewing gum for one hour prior to testing.

SUBJECT TRAINING

Participants were initially trained in the use of the general Labeled Magnitude Scale (gLMS) following the published standard procedures (Green et al., 1996; Green et al., 1993) except the top of the scale was described as the strongest imaginable sensation of any kind (Bartoshuk, 2000). The gLMS is a psychophysical tool that requires participants to rate perceived intensity along a vertical axis lined with adjectives: barely detectable = 1.5, weak = 6, moderate = 17, strong = 35, very strong = 52, strongest imaginable = 100; the adjectives are placed semi-logarithmically, based upon experimentally determined intervals to yield data equivalent to magnitude estimation (Green et al., 1996; Green et al., 1993). The scale only shows adjectives, not numbers, to the participants, but the experimenter calculates numerical data from the scale.

Participants were trained to identify each of the five taste qualities by presenting them with exemplars. Salty taste was identified as the predominant taste quality from 150mM NaCl, bitterness as the predominant quality from 0.50mM quinine-HCl, sweetness as the predominant quality from 300mM sucrose, sourness as the predominant quality from 3mM citric acid, and umami the predominant quality
from a mixture of 100mM MSG and 50mM IMP. To help subjects understand a stimulus could elicit multiple taste qualities, 300mM urea (bitter and slightly sour) and 50mM NH₄Cl (salty, bitter, and slightly sour) were employed as training stimuli. Sucrose and NaCl were presented at three concentrations (50mM, 200mM, 400mM) to ensure subjects could rank the solutions from least to most intense. All subjects were able to identify and rank taste solutions.

STIMULI AND DELIVERY

Caffeine and 6-propylthiouracil (PROP) were purchased from Sigma Chemical (St. Louis, MO) and were Sigma-ultra grade. Quinine-HCl (QHCl) was purchased from Fluka Chemika (Buchs, Switzerland).

All solutions were prepared with deionized (di) filtered water and were stored in glass bottles at 4°C-8°C and were brought to room temperature (20°C±3°C) prior to testing. Filtered di water was used as the blank stimulus and the rinsing agent in all experiments.

All testing took place in specialized sensory-testing facility comprising of seven individual computerized booths. Each subject was isolated from other subjects by vertical dividers and there was no interaction between subjects.

DETECTION THRESHOLD DETERMINATION FOR CAFFEINE AND QUININE-HCL, AND n-PROPYLTHIOURACIL

A triangle forced-choice initially-ascending procedure was used to determine detection threshold of caffeine, QHCl, and PROP for each subject. The range of concentration used is shown in Table I: caffeine concentrations were modified from ‘ISO 3972 Method of investigating sensitivity of taste’; QHCl concentrations were 0.2 log concentration steps; and PROP concentrations were 0.125 log concentration steps.
Starting at the dilution step 3, solutions (10 ml) were presented in 30 ml plastic medicine cups in groups of three. Subjects were instructed to hold the sample in their mouth for three seconds, then expectorate. Within each set of three solutions, two were water blanks and the third was the bitter compound and subjects had to identify which one was different (triangle test). The order of presentation was randomised and could have been any of three possible orders (A/blank and B/stimulus): AAB, ABA, BAA. If subjects failed to correctly identify the odd sample, the concentration was increased one step. If subjects correctly identified the sample on two occasions, the concentration was decreased one step. The level at which the sequence changed from ascending to descending or descending to ascending, was termed a reversal. Four reversals were required and the best estimate threshold for each subject was the geometric mean of the concentration where the last miss occurred and the next higher step. There was an interstimulus interval of approximately 60 sec, during which time the subject was required to rinse with di water at least 4 times. Any one session included only one bitter compound and each session could take 30 mins to complete. The detection threshold method was repeated in a separate session to check reproducibility of detection thresholds, meaning a minimum of six sessions in total for each subject.

CONSTRUCTION OF PSYCHOPHYSICAL CURVE FOR CAFFEINE, QUININE-HCL, AND n-PROPYLTHIOURACIL

The concentration ranges for constructing a psychophysical curve for the bitter stimuli are shown in Table II. For caffeine and QHCl, subjects were presented with numbered trays that contained seven randomized solutions (10 ml) of one bitter stimulus (six concentrations from the psychophysical curve and one di water control). For PROP, the only difference was solutions were presented in ascending.
concentration order, rather than randomized order (Bartoshuk, 2000). The six concentrations for each bitter stimulus ranged from below “weak” on the gLMS to maximum practical tasting limit. Each point on an individual psychophysical curve was tested at least three times.

*Stimulus delivery*

An aliquot of 10 ml of each solution (n=7) was presented in 30 ml polyethylene medicine cups (Dynarex, NY) in randomized order (except PROP see above) on a numbered tray. Subjects rinsed with *di* water at least four times over a two minute period prior to testing. Each subject tasted, and then rated each solution for sweetness, sourness, saltiness, bitterness and umami, prior to expectorating. All subjects rinsed with *di* water four times during the interstimulus interval of 90 sec. The gLMS was used as the rating method. Each sample was tasted only once per session and there were three sessions in total as a test of reliability of rating.

Psychophysical curves were constructed for the bitter compounds for each individual subject. These curves provided the opportunity to investigate perceived bitterness correlations as a function of individual sensitivities among bitter compounds at six different concentration levels and threshold concentrations. First, the intensity ratings were adjusted for bias in scale use.

**STANDARISATION OF gLMS RATINGS WITH SWEETNESS AND WEIGHT RATINGS**

The gLMS standardisation was a modified version of Delwiche *et al.*, (2001). Briefly, subjects rated the sweetness and total intensity of 10ml samples of five concentrations of sucrose (50, 100, 150, 250, 400 mM). Between each sample, subjects rinsed four times with deionized water. Subjects also rated the heaviness of
five visually identical weights (opaque, sand-filled jars at levels 52, 294, 538, 789, and 1028 g). All ratings were made on the gLMS. Subjects were asked to rate the intensity of taste or heaviness, and all judgments were made within the context of the full range of sensations experienced in life. All stimuli were presented twice in blocks of ascending order. Subjects first rated the heaviness of weights, then the intensity of sucrose solutions.

There was a significant correlation between sucrose sweetness and heaviness ratings ($r^2=0.49$, $p<0.05$). Since these sensory modalities were assumed to be unrelated, the significant correlation indicated that the gLMS ratings were prone to individual scale-use bias and required standardization across subjects.

To determine a standardization factor, each subject’s average intensity for heaviness was divided by the grand mean for heaviness across weight levels and subjects. Each individual’s bitter intensity ratings for caffeine, QHCl and PROP were multiplied by his or her personal standardization factor for scale-use bias.

STATISTICAL ANALYSIS

Data used for correlation analysis were the detection threshold concentrations and the individual bitterness intensity ratings (gLMS) at stated concentration levels. Correlation analysis (Pearson product-moment coefficients) was performed using SPSS version 12.0.1. Subjects who are termed insensitive to the bitter compounds tested have a higher detection threshold and lower intensity rating than sensitive subjects (lower detection threshold, higher intensity rating). When this data is analyzed, what is a positive correlation will have a negative sign. Therefore, in order
to assess correlations between the detection threshold concentrations and suprathreshold intensities, positive r values were converted to negative and vice versa.

PASS statistical software (2005) was used to determine the power of this study. Assuming $r = 0.35$, $n=33$ and $\alpha<0.05$, the power of the study is 0.65. Ideally a power of 0.8 should be achieved, and with $n=33$ and $\alpha<0.05$, the $r$ value = 0.45. The study was large enough to assume a type II error is within acceptable range.
Results

Detection threshold

The mean detection threshold and standard error for caffeine, QHCl and PROP was $1.2 \pm 0.12$ mM, $0.0083 \pm 0.001$ mM, and $0.088 \pm 0.07$ mM respectively. The relationship between detection thresholds for caffeine and QHCl among subjects was investigated using Pearson product-moment correlation coefficient. There was no correlations between detection thresholds for caffeine, QHCl and PROP [n=33, r=-0.006 - -0.24, p=0.97 - 0.18] (Figure 2).

Suprathreshold intensities

Psychophysical curves were constructed for caffeine, QHCl, and PROP and there was much individual variation in bitterness perception (Figure 3 A, B, & C). Even though bitterness intensity varied among subjects, as the concentration of QHCl and PROP increased there was ordinal increases in bitterness intensity across subjects, and as expected, Pearson coefficient correlations revealed a significant relationship between all points on a bitter compounds psychophysical curve [(QHCl; r=0.61-0.88, p<0.001) (PROP; r=0.65-0.924, p<0.001)]. ANOVA results showed significant differences between all incremental steps on the psychophysical curves (p<0.05). This indicates that when a subject is given increasing concentrations of quinine or PROP (above detection threshold) there is an ordinal increase in bitterness intensity relative to intensity ratings across all subjects (a subject who was insensitive to the bitter taste of the stimulus remains insensitive in relation to the other subjects for the concentrations tested). The strong correlation was also evident for caffeine, but only at the higher concentrations 12mM-72mM (r=0.61-0.96, p<0.001). Whereas, at 6mM caffeine there was a strong correlation with 3mM caffeine (r=0.63, p<0.001) and 12mM (r=0.61, p<0.001), and weaker correlations with higher caffeine concentration
Complexity of bitter taste

(r=0.43-0.46, p<0.05). The bitterness intensity ratings of the subjects at lowest concentration of caffeine (3mM) did not correlate with any of the concentrations above 6mM (r=-0.06-0.2, p=0.2-0.9). This indicates a low concentration and high concentration mechanism responsible for the perceived bitter taste of caffeine.

There were no significant correlations with subjects intensity rating of caffeine and PROP (r=-0.06 – 0.1, p=0.82 – 0.5), or quinine-HCl and PROP (r=0.07 – 0.3, p=0.72 – 0.07), which is similar to other studies investigating correlations of bitter compounds with PROP bitterness (Delwiche et al., 2001; Hansen et al., 2006; Keast et al., 2003b). Therefore sensitivity to the bitterness of PROP does not predicate that the subject will be sensitive to the bitterness of caffeine or QHCl. At the three highest concentrations of caffeine and QHCl tested, there were significant correlations (r=0.56-0.36, p<0.05). This supports previous research indicating perceptual and genetic similarities between the bitterness of caffeine and QHCl (Delwiche et al., 2001; Hansen et al., 2006)

Detection threshold and suprathreshold intensity among compounds

Table III shows Pearson product-moment correlation coefficient for detection threshold concentration and suprathreshold intensities for the individual bitter compounds across subjects. There was no significant correlation between detection threshold and suprathreshold intensity ratings for caffeine. Surprisingly there was a negative correlation between threshold of QHCl and suprathreshold intensity ratings of QHCl. This indicates that subjects who were sensitive to quinine-HCl (low threshold concentrations) generally found higher concentrations of quinine-HCl less bitter, while subjects who were insensitive to QHCl (high threshold concentrations) perceived higher concentrations of QHCl more bitter. There were positive
correlations between PROP threshold and suprathreshold intensity rating (except at the lowest concentration on the psychophysical curve).
Discussion

The relationship between the concentration of a chemical and the perception of that chemical (intensity and liking) is complex (Amerine et al., 1965; Bartoshuk, 2000; Mojet et al., 2005). The results from this study do not diminish that complexity, indeed they add to complex relationship between chemical concentration, detection threshold, and suprathreshold intensity. As the concentration of a chemical increases from detection threshold to suprathreshold there was a significant positive correlation for PROP, a significant negative correlation with QHCl, and no correlation for caffeine. The complexity may be due to multiple perceptual and peripheral mechanisms of bitter taste and these multiple mechanisms may be activated at different concentrations. Figure 4 illustrates the positive and negative correlations among chemical concentration, detection threshold, and suprathreshold intensity observed in this study. As the statistics infer, Figure 4 is a generalization of results from this study and not all subjects will follow the model.

6-n-Propylthiouracil

In this study, PROP observed the intuitive model of sensitivity throughout a concentration range with sensitivity at low concentration predicting sensitivity at higher concentrations (Figure 4). However, in a comprehensive review of variation in taste perception, Bartoshuk (2000) has previously stated relying on detection thresholds for PROP may cause misclassification of subjects ‘taster’ status in the suprathreshold range. In support of Bartoshuk’s observation, classifying PROP taster status on detection thresholds would have resulted misclassification of 4 of the 33 subjects at suprathreshold intensity, even though there was a significant correlation between detection threshold and suprathreshold sensitivity for PROP. The ability to taste PROP has been linked to the bitter receptor gene hTAS2R38 (Duffy et al.,
2004), and there is a very close association between absolute detection threshold and hTAS2R38 haplotypes (Bufe et al., 2005). As there is one known receptor linked to perception of PROP, it is not surprising to find a significant relationship between detection threshold and suprathreshold intensity. However, even for the PROP, there is speculation that additional genetic or environmental controls govern bitter taste perception as the PROP concentration increases (Bufe et al., 2005).

**Quinine-HCl**

In this study, there was a negative correlation between QHCl detection threshold concentration and suprathreshold intensity. Figure 4 illustrates a subset of the sample population have a compressed perceived intensity range relative to chemical concentration, while a second sub-set of the population have an expansive perceived intensity range relative to the chemical concentration. There have been few reports of such negative correlations between threshold and suprathreshold sensitivity within taste, although Mojet et al., (2005) reported similar negative correlations for salt and umami qualities. The psychophysical data for QHCl suggests at least two perceptual mechanisms, an independent factor regulating threshold detection, which co-varies with mechanism/s associated with suprathreshold intensities. Multiple perceptual mechanisms of QHCl is supported by multiple peripheral mechanisms, including the ability of quinine to block K+ channels (Kinnamon & Cummings, 1992), and in addition to sharing genetic factors associated with variation in perception with caffeine, QHCl has a putative specific genetic factor regulating only its bitterness perception (Bachmanov et al., 1996; Hansen et al., 2006).

**Caffeine**

Caffeine results were the most intriguing of the compounds tested. There was no correlation between detection sensitivity and sensitivity to caffeine at any point of
Complexity of bitter taste

The psychophysical function. Moreover, there was a specific concentration (6mM) where perceived bitter taste could be differentiated – the lower concentrations elicited bitterness that was correlated among subjects, the same for the higher concentrations. However, the bitterness elicited by \( \leq 6 \text{mM} \) and \( \geq 6 \text{mM} \) concentrations did not correlate with each other. Overall there were three perceptual shifts associated with caffeine concentration, which may indicate three different bitter taste mechanisms: one for detection threshold (very low concentrations, \( \leq 1 \text{mM} \)); one for \( \sim 1 - <6 \text{mM} \) concentrations of caffeine; and one for \( >6 \text{mM} \) concentrations of caffeine. Multiple perceptual mechanisms for caffeine bitterness is supported by multiple independent putative mechanisms: caffeine can translocate through cellular membranes and has the ability to interfere with 2\textsuperscript{nd} messenger systems (Peri et al., 2000); the bitterness of caffeine has been associated with the bitterness of QHCl (this study and (Delwiche et al., 2001)); and there is a proposed small (2\%) genetic link between PROP and caffeine (hTAS2R38) (Hansen et al., 2006).

Detection threshold and suprathreshold intensity among compounds

In this study there was no correlation between the detection thresholds of all three compounds, therefore sensitivity to bitter compounds at threshold level was not common across subjects. This suggests that caffeine, QHCl, and PROP have independent mechanisms responsible for their detection at low concentration. This was not surprising for PROP, as previous research has established no common bitterness with caffeine and QHCl at suprathreshold level, a result that was replicated in the present study. Previous research has shown an association between caffeine and QHCl at suprathreshold intensities (Delwiche et al., 2001; Hansen et al., 2006), a finding that was also replicated in the present study. However, at lower concentrations there was no correlation indicating the commonality in bitterness.
between caffeine and QHCl may be due to a bitter taste mechanism/s activated at higher concentrations of the two compounds.

*Organization of the bitter taste system*

If a single receptor was responsible for detection and suprathreshold intensity you would expect a strong correlation between chemical concentration, detection threshold, and suprathreshold intensity, and this was observed with PROP (Figure 4). However, if there are multiple taste transduction mechanisms that are activated at varying concentrations of the chemical there may be no association between detection threshold and suprathreshold intensity, and this was observed with caffeine. A negative association may occur if a high affinity receptor process was activated at very low concentrations of the chemical, but high enough to reach a detection threshold, then as the concentration was increased a lower affinity receptor mechanism was activated and was responsible for a perceived quality. If a subject had a larger quantity of one of the two receptor types we may expect a negative association between detection threshold and suprathreshold intensity, and this was observed with QHCl (Figure 4).

The variation and lack of correlation in bitter taste perception may be due to multiple factors. Recent advances in our knowledge of the peripheral organization of the taste system strongly indicate that taste receptor cells are quality specific (Huang *et al.*, 2006; Mueller *et al.*, 2005). In addition to this, not all bitter taste cells contain all bitter taste receptors, but sub-sets of receptors located on bitter taste cells (Chandrashekar *et al.*, 2000). Variation in receptor sub-sets of receptors on bitter taste cells may influence bitter taste perception. For example, sweet and umami taste are activated by hetero-dimers of the TAS1R family, and it is not inconceivable the
same dimer system could occur with the TAS2Rs on bitter taste cells. If a bitter taste cell lacks one part of a dimer, activation of that cell would not occur. There may also be single nucleotide polymorphisms in TAS2Rs that result in differences in bitter taste perception (Bufo et al., 2005). Moreover, each TAS2R may have multiple binding sites that are low or high affinity, and as the concentration of a compound increases the lower affinity receptor or active site of the receptor is activated (Galindo-Cuspinera et al., 2006).

Within an individual, the strength of an afferent signal may be magnified relative to other individuals. There may also be inter-individual variations in the signal processing in the human brain, although our understanding of gustatory processing in the brain is still in it’s infancy (Small, 2006).
Conclusions

There is a complex relationship between chemical concentration, detection threshold, and suprathreshold intensity of bitter compounds. The sensitivity of a person to detect very low concentrations of a compound is not necessarily associated with their sensitivity to the same compound when it is perceivably bitter. Moreover, in some situations threshold sensitivity to a compound may be inversely related to the intensity of perceived bitterness of that compound. Such complexity has practical implications as threshold determination methods are increasingly (and incorrectly) used to infer suprathreshold intensity of specific compounds e.g., Taste Dilution Analysis, (Frank et al., 2001; Ottinger et al., 2003). More broadly, this paper also continues to support that attempts to link threshold measures to food sensations and intake are at best misguided.

The bitter taste system may have distinct perceptual stages, one for threshold and at least one for suprathreshold intensities and these perceptual stages may relate to distinct oral peripheral mechanisms. As the concentration of a compound increases, receptors that have a lower affinity for the compound may become involved in the process of taste transduction, resulting in perceptual phases that can be differentiated using psychophysical methods of evaluation.
Acknowledgements

We thank Professor Sing Kai Lo for his advice on the statistics undertaken in this study. Financial support was received from the School of Exercise and Nutrition Science, Deakin University. We thank all the subjects who took part in this study.
Complexity of bitter taste

References


TABLES

Table I Concentrations and dilution steps used to determine subject detection threshold for caffeine, quinine-HCl (QHCl), and n-propylthiouracil (PROP) in water. The concentration series for caffeine was adapted from ISO3970, ‘Method of investigating sensitivity of taste’, the concentration series for quinine-HCl was prepared with successive 0.15 log dilutions with filtered deionised water, and the concentration series for PROP was prepared with successive 0.125 log dilution steps.

<table>
<thead>
<tr>
<th>Caffeine [mM]</th>
<th>Quinine-HCl [mM]</th>
<th>PROP [mM]</th>
<th>Dilution step</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.28</td>
<td>0.00064</td>
<td>0.01</td>
<td>1</td>
</tr>
<tr>
<td>0.33</td>
<td>0.0009</td>
<td>0.014</td>
<td>2</td>
</tr>
<tr>
<td>0.42</td>
<td>0.0013</td>
<td>0.019</td>
<td>3</td>
</tr>
<tr>
<td>0.52</td>
<td>0.0017</td>
<td>0.025</td>
<td>4</td>
</tr>
<tr>
<td>0.66</td>
<td>0.0025</td>
<td>0.033</td>
<td>5</td>
</tr>
<tr>
<td>0.80</td>
<td>0.0035</td>
<td>0.045</td>
<td>6</td>
</tr>
<tr>
<td>1.03</td>
<td>0.005</td>
<td>0.059</td>
<td>7</td>
</tr>
<tr>
<td>1.3</td>
<td>0.007</td>
<td>0.079</td>
<td>8</td>
</tr>
<tr>
<td>1.57</td>
<td>0.01</td>
<td>0.1</td>
<td>9</td>
</tr>
<tr>
<td>1.84</td>
<td>0.014</td>
<td>0.14</td>
<td>10</td>
</tr>
<tr>
<td>2.11</td>
<td>0.02</td>
<td>0.19</td>
<td>11</td>
</tr>
<tr>
<td>2.38</td>
<td>0.028</td>
<td>0.25</td>
<td>12</td>
</tr>
<tr>
<td>2.65</td>
<td>0.04</td>
<td>0.33</td>
<td>13</td>
</tr>
</tbody>
</table>
Table II Concentrations of caffeine, quinine-HCl (QHCl), and n-propylthiouracil (PROP) used to generate psychophysical curves

<table>
<thead>
<tr>
<th>Caffeine [mM]</th>
<th>QHCl [mM]</th>
<th>PROP [mM]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>0.05</td>
<td>0.05</td>
</tr>
<tr>
<td>6</td>
<td>0.1</td>
<td>0.25</td>
</tr>
<tr>
<td>12</td>
<td>0.15</td>
<td>0.75</td>
</tr>
<tr>
<td>24</td>
<td>0.2</td>
<td>1.25</td>
</tr>
<tr>
<td>48</td>
<td>0.25</td>
<td>2.5</td>
</tr>
<tr>
<td>72</td>
<td>0.3</td>
<td>5.5</td>
</tr>
</tbody>
</table>
Table III  Pearsons product-moment correlation between threshold and six suprathreshold intensity ratings for caffeine, quinine-HCl (QHCl), and 6-n-propylthiouracil (PROP). Concentrations of chemicals for stimulus # 1-6 are shown in Table II.  ns denotes not significant, * denotes significant at p<0.05, ** denote significance p<0.01

<table>
<thead>
<tr>
<th>Stimulus #</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caffeine</td>
<td>0.001 ns</td>
<td>0.15 ns</td>
<td>-0.2 ns</td>
<td>-0.09 ns</td>
<td>-0.08 ns</td>
<td>-0.05 ns</td>
</tr>
<tr>
<td>QHCl</td>
<td>-0.08 ns</td>
<td>-0.38*</td>
<td>-0.4*</td>
<td>-0.36*</td>
<td>-0.37*</td>
<td>-0.33*</td>
</tr>
<tr>
<td>PROP</td>
<td>0.26 ns</td>
<td>0.43**</td>
<td>0.43**</td>
<td>0.46**</td>
<td>0.4*</td>
<td>0.43**</td>
</tr>
</tbody>
</table>
FIGURES

**Figure 1** Schematic illustration of the relationship between chemical concentration, detection threshold, and suprathreshold intensity using general Labeled Magnitude Scale (gLMS). The left hand side of the bold black Y-axis represents chemical concentration from 0 molar (0 M) solution to a saturated solution. The right hand side of the bold black Y-axis represents the perceptual relationship to increasing concentration. The far right vertical axis represents the gLMS scale from no perception to a theoretical terminal threshold.

**Figure 2** Detection threshold correlation. Detection threshold concentrations for caffeine, quinine-HCl (QHCl) and n-propylthiouracil (PROP) on a three-dimensional plot. All concentrations are in mM, the y-axis is caffeine, x-axis is QHCl, and the z-axis PROP. Each point represents the threshold concentrations for 1 of the 33 subjects.

**Figure 3 A-C** Psychophysical curves of the sample population mean and examples of an insensitive and sensitive subject for A/ caffeine, B/ quinine-HCl, and C/ PROP. Included in each graph is a sensitive (highest curve) and insensitive subject (lowest curve) for that compound as well as the mean psychophysical curve. The Y-axis is a numerical measure of bitterness intensity from the general Labelled Magnitude Scale (gLMS). The x-axis has two labels, the upper label in the log millimolar concentration for the particular compound, the lower label is the actual millimolar concentration. Error bars represent standard errors.

**Figure 4** Schematic illustration of the association between chemical concentration, detection threshold, and suprathreshold intensity for propylthiouracil (PROP) and quinine-HCl (QHCl). The bold black solid vertical line represents the chemical concentration. The thin solid vertical lines represent the gLMS intensity
rating relative to the chemical concentration. The bottom of each thin solid line (—) represents the detection threshold. The top of each solid line represents an intensity of ~20 on the gLMS scale. The vertical dashed line below the solid line represents the concentrations of chemical in solution without eliciting a noticeable difference from water. The left hand side of the chemical concentration axis illustrates results observed for PROP, with sensitivity at detection threshold consistent over the concentration range tested. This is illustrated by an equal perceived intensity range relative to chemical concentration regardless of an individual's sensitivity to PROP. The right hand side of the concentration axis illustrates the results observed for QHCl with subjects rating between 0-20 gLMS as either a compressed perceived intensity range relative to chemical concentration, or an expansive perceived intensity range (far right) relative to the chemical concentration.
Complexity of bitter taste

**Concentration continuum**

- Saturated solution
- Terminal threshold
- Detection threshold: Chemical in solution but no perception

**Perceptual phases**

- Recognition threshold
- Dynamic phase

- ~20 gLMS
- 0 gLMS

Perceived intensity region
Detection threshold correlation
Caffeine

![Graph showing the relationship between log caffeine concentration and GLMS intensity for sensitive and insensitive subjects.](image)

- **Group mean bitterness**
- **Sensitive subject**
- **Insensitive subject**

<table>
<thead>
<tr>
<th>[log caff mM]</th>
<th>0.4</th>
<th>0.6</th>
<th>0.8</th>
<th>1.0</th>
<th>1.2</th>
<th>1.4</th>
<th>1.6</th>
<th>1.8</th>
<th>2.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>[caffeine mM]</td>
<td>3</td>
<td>6</td>
<td>12</td>
<td>24</td>
<td>48</td>
<td>72</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Complexity of bitter taste

QHCl

![Graph showing the relationship between [log QHCl] and gLMS intensity for group mean bitterness, sensitive subject, and insensitive subject. The x-axis represents [log QHCl] with values from -1.4 to -0.4, and the y-axis represents gLMS intensity with values from 0 to 40. The graph includes data points and error bars for each condition.]
Complexity of bitter taste

PROP

\[ \log(\text{PROP}) \]

\begin{array}{cccccc}
-1.5 & -1.0 & -0.5 & 0.0 & 0.5 & 1.0 \\
\hline
0 & 10 & 20 & 30 & 40 & 50 \\
\end{array}

Group mean bitterness
Sensitive subject
Insensitive subject

\begin{array}{cccccccc}
0.05 & 0.25 & 0.75 & 1.25 & 2.5 & 5.5 \\
\end{array}
Complexity of bitter taste

Fig 4

PROP Model (Intuitive)

- Insensitive
- Sensitivity threshold intensity ~20 gLMS
- Detection threshold 0 gLMS
- Equal perceived intensity in relation to chemical concentration

QHCl Model

- Insensitive
- Sensitive ~20 gLMS
- Chemical in solution but no perception
- Compressive
- Expansive

Chemical Concentration axis

Insensitive
Sensitive
0 M

Supra threshold intensity

Chemical in solution but no perception