The cellular and molecular basis of bitter tastant-induced bronchodilation

Cheng-Hai Zhang, University of Massachusetts Medical School

Let us know how access to this document benefits you.
Follow this and additional works at: https://escholarship.umassmed.edu/faculty_pubs

Part of the Amino Acids, Peptides, and Proteins Commons, Animal Experimentation and Research Commons, Cells Commons, Cellular and Molecular Physiology Commons, Chemical Actions and Uses Commons, Investigative Techniques Commons, Molecular Biology Commons, Pharmaceutical Preparations Commons, Respiratory System Commons, Respiratory Tract Diseases Commons, and the Therapeutics Commons

Repository Citation
Zhang C, Lifshitz LM, Uy K, Ikebe M, Fogarty KE, ZhuGe R. (2013). The cellular and molecular basis of bitter tastant-induced bronchodilation. University of Massachusetts Medical School Faculty Publications. https://doi.org/10.1371/journal.pbio.1001501. Retrieved from https://escholarship.umassmed.edu/faculty_pubs/247

This material is brought to you by eScholarship@UMassChan. It has been accepted for inclusion in University of Massachusetts Medical School Faculty Publications by an authorized administrator of eScholarship@UMassChan. For more information, please contact Lisa.Palmer@umassmed.edu.
The Cellular and Molecular Basis of Bitter Tastant-Induced Bronchodilation

Cheng-Hai Zhang¹*, Lawrence M. Lifshitz²³, Karl F. Uy⁴, Mitsuho Ikebe¹, Kevin E. Fogarty²³, Ronghua ZhuGe¹²*

1 Department of Microbiology and Physiological Systems, University of Massachusetts Medical School, Worcester, Massachusetts, United States of America, 2 Biomedical Imaging Group, University of Massachusetts Medical School, Worcester, Massachusetts, United States of America, 3 Program in Molecular Medicine, University of Massachusetts Medical School, Worcester, Massachusetts, United States of America, 4 Department of Surgery, Division of Thoracic Surgery, University of Massachusetts Memorial Medical Center, Worcester, Massachusetts, United States of America

Abstract

Bronchodilators are a standard medicine for treating airway obstructive diseases, and β2 adrenergic receptor agonists have been the most commonly used bronchodilators since their discovery. Strikingly, activation of G-protein-coupled bitter taste receptors (TAS2Rs) in airway smooth muscle (ASM) causes a stronger bronchodilation in vitro and in vivo than β2 agonists, implying that new and better bronchodilators could be developed. A critical step towards realizing this potential is to understand the mechanisms underlying this bronchodilation, which remain ill-defined. An influential hypothesis argues that bitter tastants generate localized Ca²⁺ signals, as revealed in cultured ASM cells, to activate large-conductance Ca²⁺-activated K⁺ channels, which in turn hyperpolarize the membrane, leading to relaxation. Here we report that in mouse primary ASM cells bitter tastants neither evoke localized Ca²⁺ events nor alter spontaneous local Ca²⁺ transients. Interestingly, they increase global intracellular [Ca²⁺]i, although to a much lower level than bronchoconstrictors. We show that these Ca²⁺ changes in cells at rest are mediated via activation of the canonical bitter taste signaling cascade (i.e., TAS2R-gustducin-phospholipase Cβ [PLCβ]-inositol 1,4,5-triphosphate receptor [IP3R]), and are not sufficient to impact airway contractility. But activation of TAS2Rs fully reverses the increase in [Ca²⁺]i induced by bronchoconstrictors, and this lowering of the [Ca²⁺]i is necessary for bitter tastant-induced ASM cell relaxation. We further show that bitter tastants inhibit L-type voltage-dependent Ca²⁺ channels (VDCCs), resulting in reversal in [Ca²⁺]i, and this inhibition can be prevented by pertussis toxin and G-protein βγ subunit inhibitors, but not by the blockers of PLCβ and IP3R. Together, we suggest that TAS2R stimulation activates two opposing Ca²⁺ signaling pathways via Gβγ to increase [Ca²⁺], at rest while blocking activated L-type VDCCs to induce bronchodilation of contracted ASM. We propose that the large decrease in [Ca²⁺], caused by effective tastant bronchodilators provides an efficient cell-based screening method for identifying potent dilators from among the many thousands of available bitter tastants.

Citation: Zhang C-H, Lifshitz LM, Uy KF, Ikebe M, Fogarty KE, et al. (2013) The Cellular and Molecular Basis of Bitter Tastant-Induced Bronchodilation. PLoS Biol 11(3): e1001501. doi:10.1371/journal.pbio.1001501

Introduction

Airway obstructive diseases (asthma and chronic obstructive pulmonary disease [COPD]) have become increasingly prevalent, currently affecting more than 300 million people worldwide. Dysfunction of airway smooth muscle (ASM) cells, a major cell type in the respiratory tree, plays a pivotal role in promoting progression of these diseases and in contributing to their symptoms of these diseases [1–3]. With their ability to contract and relax, these cells regulate the diameter and length of conducting airways, controlling dead space and resistance to airflow to and from gas-exchanging areas. Their excessive contraction, as seen in patients with asthma and COPD, can fully close the airways, thereby preventing gas exchange and threatening life. Not surprisingly, bronchodilators have been used as the medication of choice for asthmatic attacks and as a standard medicine for managing COPD [4,5]. However, available bronchodilators have adverse side effects, and are not sufficiently effective for severe asthmatics and many other COPD patients. A better understanding of the mechanisms regulating ASM thus holds the promise of developing more effective and safe bronchodilators, which in turn would have a significant impact in reducing mortality and morbidity caused by asthma and COPD.

Bitter tastants represent a new class of compounds with potential as potent bronchodilators. Deshpande et al. recently found that cultured ASM cells express G-protein-coupled bitter taste receptors (TAS2Rs) [6], a class of proteins long thought to be expressed only in the specialized epithelial cells in the taste buds of...
Author Summary

Bitter taste receptors (TAS2Rs), a G-protein-coupled receptor family long thought to be solely expressed in taste buds on the tongue, have recently been detected in airways. Bitter substances can activate TAS2Rs in airway smooth muscle to cause greater bronchodilation than β2 adrenergic receptor agonists, the most commonly used bronchodilators. However, the mechanisms underlying this bronchodilation remain elusive. Here we show that, in resting primary airway smooth muscle cells, bitter tastants activate a TAS2R-dependent signaling pathway that results in an increase in intracellular calcium levels, albeit to a level much lower than that produced by bronchoconstrictors. In bronchoconstricted cells, however, bitter tastants reverse the bronchoconstrictr-induced increase in calcium levels, which leads to the relaxation of smooth muscle cells. We find that this reversal is due to inhibition of L-type calcium channels. Our results suggest that under normal conditions, bitter tastants can activate TAS2Rs to modestly increase calcium levels, but that when smooth muscle cells are constricted, they can block L-type calcium channels to induce bronchodilation. We postulate that this novel mechanism could operate in other extraoral cells expressing TAS2Rs.

the tongue that allow organisms to avoid harmful toxins and noxious substances characterized by bitterness [7–10]. Importantly, bitter tastants with diverse chemical structures cause greater ASM relaxation in vitro than β2 adrenergic agonists, the most commonly used bronchodilators to treat asthma and COPD [6,11]. Moreover, these compounds can effectively relieve in vivo asthmatic airway obstruction than β2 adrenergic agonists in a mouse model of asthma [6], making them highly attractive bronchodilators for asthma and COPD.

Bitter tastant-induced bronchodilation was unexpected, because these agents appeared to increase intracellular Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]\(_i\)) to a level comparable to that produced by potent bronchoconstrictors [6], which should have led to smooth muscle contraction [12]. To reconcile this apparent paradox, it was proposed that bitter tastants activate the canonical bitter taste signaling pathway (i.e., TAS2R-gustducin-phospholipase Cβ [PLCβ]-inositol 1,4,5-triphosphate receptor [IP3R]) to increase focal Ca\(^{2+}\) release from endoplasmic reticulum, which then activate large-conductance Ca\(^{2+}\)-activated K\(^+\) channels thereby hyperpolarizing the membrane [6]. However, we subsequently demonstrated through patch-clamp recordings that bitter tastants do not activate large-conductance Ca\(^{2+}\)-activated K\(^+\) channels but rather inhibit them [11]. Moreover, three different large-conductance Ca\(^{2+}\)-activated K\(^+\) channel blockers did not affect the bronchodilation induced by bitter tastants [11]. Therefore, a different mechanism must be responsible for the bitter tastant-induced bronchodilation.

The apparent conundrum of putative [Ca\(^{2+}\)]\(_i\) elevation leading to relaxation may be attributed to the fact that Ca\(^{2+}\) responses to bitter tastants were assessed in cultured human ASM cells, while the contractile responses to them were investigated in freshly dissected ASM tissues [6]. It is well known that cultured smooth muscle cells lines alter their phenotype (i.e., losing their ability to contract and relax [13,14]) and it is likely their Ca\(^{2+}\) response is also modified. Therefore, to understand bitter tastant-induced bronchodilation, it is necessary to study the contraction and the underlying signaling in freshly isolated ASM tissues and cells. Using this approach in the present study, we found that bitter tastants activate the canonical bitter taste signaling cascade, slightly increasing global [Ca\(^{2+}\)]\(_i\) in resting cells, but not to a level sufficient to cause contraction. However, bitter tastants reverse the increase in [Ca\(^{2+}\)]\(_i\), evoked by bronchoconstrictors, leading to bronchodilation. This reversal is mediated by the suppression of L-type voltage-dependent Ca\(^{2+}\) channels (VDCCs) in a gustducin βγ subunit-dependent, yet PLCβ- and IP3R-independent manner. Hence, we propose that TAS2R activation in ASM stimulates two opposing Ca\(^{2+}\) signaling pathways, both mediated by Gβγ subunits, which increases [Ca\(^{2+}\)]\(_i\) at rest but blocks activated L-type VDCCs reversing the contraction they cause. These results provide the cellular and molecular basis of bitter tastant-induced bronchodilation that occurs in vitro and in vivo. They further reveal a Ca\(^{2+}\) signal that is well suited for screening and identifying potent bronchodilators from among the many thousands of available bitter tastants.

Results

To uncover the mechanism underlying bitter tastant-induced bronchodilation as demonstrated in both in vitro and in vivo normal and asthmatic models of mice, and in vitro human airways ([6,11,15,16], we examined how bitter tastants affected both [Ca\(^{2+}\)]\(_i\), and ASM contraction in freshly isolated airway cells and tissues from mouse and human. Fluo-3 was used to assess the effect of bitter tastants on [Ca\(^{2+}\)]\(_i\); chloroquine and denatonium, two substances commonly used to study bitter taste signaling, were used as bitter tastants.

Bitter Tastants Modestly Raise Global [Ca\(^{2+}\)]\(_i\), with No Change in Force Generation in Native ASM at Rest

We started our analysis by examining the Ca\(^{2+}\) response to bitter tastants in resting cells. In contrast to the marked increase in global [Ca\(^{2+}\)]\(_i\), reported in resting cultured human ASM cells [6], we observed, in resting native ASM cells from mouse, that chloroquine (0.1 μM–1 mM) only modestly raised global [Ca\(^{2+}\)]\(_i\), to a level much lower than when cells contracted after application of Mch at 0.1 μM–100 μM (Figure 1A and Figure S1A). Chloroquine (330 μM) increased fluo-3 fluorescence (ΔF/ F\(_0\) (i.e., [Ca\(^{2+}\)]\(_i\)) both in the presence of extracellular Ca\(^{2+}\) (37.7%±8%, n = 19) and in its absence (29.3%±6%, n = 15; p>0.05), indicating that the source for this chloroquine response is from internal Ca\(^{2+}\) stores.

To examine whether this modest increase in [Ca\(^{2+}\)]\(_i\)), is sufficient to trigger contraction, we measured smooth muscle force formation in mouse airways. As shown in (Figure 1B and S1B), chloroquine (10 μM–1 mM) did not cause contraction of mouse airways, although there was a tendency to decrease the basal tone of airways. As a comparison, Mch at concentrations between 0.3 μM and 10 μM induced contraction markedly and in a dose-dependent manner (Figure 1B and S1B).

Bitter Tastants Do Not Generate Localized Ca\(^{2+}\) Events

Mouse ASM cells exhibit spontaneous Ca\(^{2+}\) sparks resulting from the opening of ryanodine receptors in the sarcoplasmic reticulum [17]. To test whether bitter tastants generate local Ca\(^{2+}\) events as proposed by Deshpande et al. [6], we stimulated ASM cells with chloroquine (10 μM, a concentration around EC50) for 2 min and measured Ca\(^{2+}\) sparks. Off 40 chloroquine-stimulated cells, 27 cells generated a global [Ca\(^{2+}\)]\(_i\), increase that precluded an accurate estimate of Ca\(^{2+}\) sparks. In the remaining 13 cells without a detectable global rise in [Ca\(^{2+}\)]\(_i\), chloroquine inhibited the spark frequency but had no effect on the amplitude (frequency [Hz]: 2.13±0.24 in control and 1.62±0.21 with chloroquine [n = 13,
p<0.05, paired Student’s t-test]; amplitude (ΔF/F₀ at the brightest location): 20.6±1.69 in control and 18.1±1.3 with chloroquine (n=13, p>0.05, paired Student’s t-test]). To test whether spontaneous Ca²⁺ sparks mask the effect of bitter tastants on other forms of local Ca²⁺ releases, such as Ca²⁺ puffs due to the opening of IP3Rs [18], we examined the Ca²⁺ responses to chloroquine in ASM cells pretreated with 100 μM ryanodine. In these cells, prior to chloroquine application, no spontaneous sparks were observed (n=14). Chloroquine (10 μM) increased global [Ca²⁺] by 12%±4% (ΔF/F₀ at its brightest location) in nine cells, and failed to cause any detectable Ca²⁺ increase in five cells. There were no detectable local Ca²⁺ events produced in any of the 14 cells. These results indicate that chloroquine at 10 μM does not increase local Ca²⁺ events (either Ca²⁺ puffs or Ca²⁺ sparks).

Bitter Tastants Activate the TAS2R Signaling Pathway to Modestly Raise Global [Ca²⁺] in Native ASM at Rest

We next examined the cause of the modest global [Ca²⁺] rise by bitter tastants. Since in taste cells, bitter tastants bind to TAS2R to activate the pertussis toxin (PTX) sensitive G-protein gustducin, which in turn induces a PLCβ2 and IP3 signaling cascade [19,20], we studied whether bitter tastants activate this TAS2R signaling pathway. In native ASM cells, PTX (1 μg/ml, and 6–8 h pretreatment), reduced the chloroquine-induced increase in global

Figure 1. Bitter tastants modestly increase intracellular Ca²⁺ concentration ([Ca²⁺]i) by activating a canonical TAS2R signaling cascade. (A) Chloroquine (Chloro) raised [Ca²⁺]i to a level much less than Mch. [Ca²⁺]i was measured with fluo-3 in the form of acetoxymethyl ester, loaded into isolated mouse ASM cells, and expressed as ΔF/F₀ (%). (B) 1 mM chloro did not contract airways (using tension as its proxy) while 100 μM Mch caused a robust contraction. Data are mean ± SEM (n=6 for chloro, and n=5 for Mch). (C) PTX, gallein, anti-βγ (MPS-phanosducin-like protein C terminus, a Gβγ blocking peptide), U73122, and 2-APB inhibited chloro-induced increase in [Ca²⁺]i (n=19–24 cells). Isolated mouse ASM cells were either pretreated with 1 μg/ml PTX for 6–8 h or with 1 μM anti-βγ for 1–2 h or with each of the other compounds listed for 5–10 min. The effects of PTX and anti-βγ were calculated by normalizing the response of chloro to that from the time matched cells without the pretreatments, and the effects of other three compounds were analyzed by normalizing the response of chloro to its own control without the compound. (D) RT-PCR transcripts after amplification with primers to TAS2R107, α-gustducin, Gβ3, Gγ13, PLCβ2, and β-actin. Note that no transcript was detected with TAS2R108 primers. RNAs were isolated from mouse tracheas and mainstem bronchi, and reactions without complementary DNA were used as a negative control. (E) Cellular distribution of TAS2R107 in three focus planes (bottom, middle, and top) of an isolated mouse ASM cell. The TAS2R107 immunostaining intensity after 3D deconvolution (see Methods) was pseudocolored with the color map on the right. This makes positive (but dim) pixels more easily distinguished from background. Eight cells showed a similar subcellular distribution pattern.

doi:10.1371/journal.pbio.1001501.g001
[Ca\(^{2+}\)]

To 21.1% ± 8.6% of the control cells (n = 20; Figure 1C).

Also both gallein (20 μM and 30 min pretreatment), a blocker of

the Gβγ dimer of PTX sensitive G proteins, and MPS-phloretin-like protein C terminus, a Gβγ blocking peptide (anti-βγ; 1 μM, and 1 h pretreatment) [21,22] reduced the bitter tansant-mediated increase in [Ca\(^{2+}\)]

rise in [Ca\(^{2+}\)]

h pretreatment) [21,22] reduced the bitter tansant-mediated increase in [Ca\(^{2+}\)], to 19.9% ± 8.5% (n = 19; Figure 1C) and 18.4% ± 4.8% of the controls, respectively. Finally, U73122

(3 μM), a blocker of PLCβ, and 2-aminoethoxydiphenyl borate

(2-APB) (50 μM), an IP3R antagonist, suppressed the bitter tansant-induced increases in [Ca\(^{2+}\)], to 18.0% ± 5.5% (n = 24) and −10.5% ± 7.3% of controls, respectively (Figure 1C). These results indicate that bitter tansants do activate the TAS2R signaling transduction pathway (i.e., TAS2R-PTX-sensitive G protein-PLCb-IP3R) to release Ca\(^{2+}\) from internal stores. This conclusion is further supported by the finding that mouse ASM cells express transcripts for TAS2R107, α-gustducin, Gβ3, Gγ13, and PLCβ2 (Figure 1D), and display peripheral localization of TAS2R107 (Figure 1E).

Bitter Tansant-Induced Bronchodilation Is Due to Reversal of the Rise in Global [Ca\(^{2+}\)] Caused by Bronchoconstrictors

Bitter tansants at μM levels can modestly increase [Ca\(^{2+}\)], in resting cells, but this raises a conundrum as they also can fully relax airways precontracted by bronchoconstrictors [6,11]. In light of the fact that an increase in [Ca\(^{2+}\)], is the primary signal for contraction in all smooth muscle, we explored how bitter tansants affect [Ca\(^{2+}\)], evoked by bronchoconstrictors. To better quantify these effects, we measured ASM Ca\(^{2+}\) response and cell shortening at the same time. The cells were stimulated with methacholine (Mch), a stable analogue of acetylcholine, which is the major neurotransmitter in parasympathetic nerves. As expected, Mch (100 μM) rapidly increased [Ca\(^{2+}\)], as fluo-3 fluorescence increased by 162% ± 26% (ΔF/F₀), and concurrently caused cell shortening by 49% ± 8% (n = 21; Figure 2A and 2B). Strikingly, chloroquine (1 mM) almost completely reversed this [Ca\(^{2+}\)], increase (i.e., bringing [Ca\(^{2+}\)], down to a level only 15% ± 2% higher than pre-stimulation levels, n = 12, p < 0.01 Mch versus Mch+chloroquine). The reversal of the increase in [Ca\(^{2+}\)], was closely associated with relaxation in ASM cells from both mouse (back to 89% ± 7% of the pre-stimulation length; Figure 2B and Video S1) and human (back to 94% ± 5% of the control length; Figure S2A). Denatonium (1 mM) generated similar effects on [Ca\(^{2+}\)], and cell shortening in response to Mch in mouse ASM cells (n = 9).

The inverse relationship between changes in [Ca\(^{2+}\)] and the resulting cell length (i.e., lowering [Ca\(^{2+}\)], results in cell lengthening) in response to bitter tansants suggests that bitter tansants reduce [Ca\(^{2+}\)], leading to bronchodilation. If this is the case, one would expect that bitter tansant-induced bronchodilation could be prevented if [Ca\(^{2+}\)], was clamped to a physiologically high level. To test this possibility, we used staphylococcal α-toxin (16,000 units/ml) to make the ASM membrane permeable to ions such that the intracellular [Ca\(^{2+}\)], could be controlled at will. A major advantage of using this toxin is that it does not damage the cells; thus signaling processes such as the G-protein-coupled receptor mediated signaling remain intact [23]. As shown in Figure 3A, raising [Ca\(^{2+}\)], to 3 μM caused a robust increase in tension in mouse airway. More importantly, at this fixed [Ca\(^{2+}\)] level, denatonium, chloroquine and quinine (all at 1 mM) failed to relax ASM in the time frame they would have in Mch contracted airways without α-toxin treatment. Therefore, clamping [Ca\(^{2+}\)], at μM levels can prevent bitter tansant-induced bronchodilation, strongly arguing that reduction of [Ca\(^{2+}\)], by bitter tansants is necessary for their relaxation action. These results further imply that a decrease in Ca\(^{2+}\) sensitivity is probably not a major mechanism underlying bitter tansant-induced bronchodilation.

The Prominent Role of L-type Ca\(^{2+}\) Channels in Mediating Mch-Induced Contraction

Having established that the suppression of [Ca\(^{2+}\)], is necessary for bitter tansant-induced bronchodilation, we next addressed the molecule(s) that bitter tansants act on to reduce [Ca\(^{2+}\)]. Before addressing this critical question, it is appropriate to determine the Ca\(^{2+}\) pathways underlying Mch-induced contraction because a controversy persists [24]. Mch activates both the M3 muscarinic acetylcholine receptor [M3R]-Gq-PLCβ-IP3 pathway and the M2 muscarinic acetylcholine receptor [M2R]-Gi/o pathway to raise

Figure 2. Bitter tansants reverse Mch-induced increase in [Ca\(^{2+}\)], and cell shortening in mouse ASM. (A) Time course of the effect of chloro (1 mM) on a 100 μM Mch-induced increase in [Ca\(^{2+}\)], (represented as ΔF/F₀ integrated over the entire cell) and cell shortening. Images show the changes in [Ca\(^{2+}\)], displayed as fluorescence intensity (rather than ΔF/F₀ to aid visualization). Cell length is indicated by the red lines. Images were taken at the time indicated on the time course of [Ca\(^{2+}\)], (upper panel), (B) Relationships between [Ca\(^{2+}\)], (left axis, blue bars) and cell length (right axis, red bars) in response to Mch and Chloro. The letters correspond to the time shown in the upper panel in (A) (n = 23 cells, mean ± SEM, *p < 0.05, **p < 0.01 using two-tailed Student’s t-test). ΔF/F₀ is zero by definition at α, so no blue bar is present at α.
by releasing Ca\(^{2+}\) from internal stores and inducing Ca\(^{2+}\) influx from the extracellular space respectively [25,26]. It has also been suggested that Ca\(^{2+}\) release from the internal stores contributes to the early phase of Mch-induced contraction, and Ca\(^{2+}\) influx is required to sustain elevated [Ca\(^{2+}\)]\(i\) and contraction. Indeed, we found that the sustained contraction by Mch in mouse ASM is largely dependent on Ca\(^{2+}\) influx (Figure S3). However, the route of Ca\(^{2+}\) influx upon muscarinic receptor activation among different species is highly debatable [24]. Many studies suggest that L-type Ca\(^{2+}\) channels are the major path of the Ca\(^{2+}\) influx for contraction [27–32]. To determine the role of this channel in mediating Mch-induced contraction in mouse airways, we examined whether L-type channel specific blockers inhibit Mch-induced contraction and the rise in [Ca\(^{2+}\)]\(i\). Previous studies showed that diltiazem, a well-known L-type Ca\(^{2+}\) channel blocker that belongs to the benzothiazepine class, dose-dependently reverses Mch-induced bronchodilation in mouse precontracted airways [31–33]. We therefore examined whether this blocker produces similar inhibition of Mch-induced airway force generation in mouse and human airways. Figures S2B and S4A show that diltiazem dose-dependently inhibited Mch-induced bronchodilation of Mch precontracted mouse airways. (A) Clamping [Ca\(^{2+}\)]\(i\), prevented bitter tasters from causing bronchodilation. Mouse airway rings were permeabilized with α-toxin (see Method), and extracellular [Ca\(^{2+}\)]\(i\) was set at 1 nM and then switched to 3 μM as indicated under the trace. Seven individual experiments (two for denatonium, two quinine, and three chloro all at 1 mM) show responses similar to that shown on the left, so the results were pooled and displayed on the right. Each ring’s normalized tension is its tension at the experiment’s end divided by its tension just prior to application of bitter tasters, times 100. (B) FPL 64176 (FPL), an L-type VDCC agonist, prevented chloro from reversing the [Ca\(^{2+}\)]\(i\) rise induced by 100 μM Mch. Left panel: a typical time course; ΔF/F\(_0\) for each curve is scaled to have a value of 100 at the peak before chloro is added. Right panel: average results of 16 cells. The values are represented as (ΔF/F\(_0\) at the peak after Mch–ΔF/F\(_0\) at 30 s after chloro)/(ΔF/F\(_0\) at the peak after Mch–ΔF/F\(_0\) at basal)×100 (i.e., the decrease due to chloro divided by the increase due to Mch). *\(p\)<0.01, control versus +FPL. (C) FPL dose-dependently reversed chloro-induced bronchodilation (using tension as a proxy measure) in Mch precontracted airways (n = 5–7 independent experiments). Data on the right panels are mean ± SEM. % relaxation = tension decrease due to chloro divided by tension increase due to Mch, ×100. The tension decrease at each concentration of chloro is measured once the tension stabilizes. The tension decrease at each increased concentration is always measured relative to the peak tension (i.e., it is total decrease, not the incremental decrease due to the additional chloro which was added).

doi:10.1371/journal.pbio.1001501.g003

Figure 3. Suppression of [Ca\(^{2+}\)]\(i\), by inhibiting L-type VDCCs is necessary for bitter tastant-induced bronchodilation of Mch precontracted mouse airways.
Moreover, it reversed the Mch-induced increase in \([\text{Ca}^{2+}]\), by 90.2\%±2.9\% in single isolated mouse ASM cells (n=12 cells). To further examine whether diltiazem inhibits muscarinic receptor-mediated contraction, we treated the airways with diltiazem at different concentrations before Mch administration. Since a unique feature of diltiazem in inhibiting L-type \text{Ca}^{2+} \text{ channels is its use-dependence of action (i.e., it more likely binds to, and therefore blocks, channels as they open), in this series of experiments we challenged the airways twice with KCl to activate the \text{Ca}^{2+} \text{ channels. As shown in Figure S4B, under this condition, diltiazem dose-dependently suppressed Mch-induced contraction and at 100 \mu\text{M} it inhibited the force by 85\%±6\% (n=5). To directly demonstrate that diltiazem inhibits L-type \text{Ca}^{2+} \text{ channels, we studied the effect of diltiazem on L-type \text{Ca}^{2+} \text{ currents with patch-clamp recording. We found that diltiazem dose-dependently reduced depolarization-induced L-type \text{Ca}^{2+} \text{ currents at concentrations over the same range as that which blocked contraction (Figure S4C). Dihydropyridines (e.g., nisoldipine, nifedipine, and isradipine) are another well-known class of L-type \text{Ca}^{2+} \text{ channel blockers [34–36]. Previously we found that dihydropyridines can inhibit L-type \text{Ca}^{2+} \text{ current in mouse ASM [17]. To further establish the role of this class of \text{Ca}^{2+} \text{ channels in mediating Mch-induced contraction, we assessed the effect of nisoldipine on the contraction evoked by Mch. It is known that dihydropyridines bind stronger to inactivated \text{Ca}^{2+} \text{ channels, thus displaying a so-called voltage dependent inhibition [37]. Therefore, in this series of experiments, to facilitate the interaction between nisoldipine and L-type \text{Ca}^{2+} \text{ channels, we examined the effect of nisoldipine on Mch-induced contraction in the presence of 20 \text{mM} KCl to modestly depolarize the membrane. Under this condition, 1 \mu\text{M} nisoldipine inhibited Mch-induced contraction by 74\%±8\% (n=5; Figure S4D). This result is similar to the inhibition of this blocker on carpachol-induced contraction in rat ASM, when it is similarly depolarized [29]. Hence, with specific L-type \text{Ca}^{2+} \text{ channel blockers of distinct structures we have established that L-type \text{VDCCs are the major contributor to \text{Ca}^{2+} \text{ influx and sustained contraction in response to Mch in mouse airways.}

**Bitter Tastants Inhibit L-Type \text{VDCCs to Decrease \text{Ca}^{2+}, Evoked by Bronchoconstrictors**

Given the prominent role of L-type \text{VDCCs in Mch-induced sustained contraction in mouse airways, and our and others' findings that bitter tastants reverse Mch-induced sustained contractile [6,11], we hypothesized that bitter tastants inhibit L-type \text{VDCCs, leading to relaxation of airways precontracted by Mch. To test this possibility, we investigated whether the L-type \text{VDCC agonist FPL 64176 [38,39] can prevent the inhibitory effect of bitter tastants on the Mch-induced \([\text{Ca}^{2+}]), rise and contraction. At the single cell level, 10 \mu\text{M} FPL prevented chloroquine from reducing the \([\text{Ca}^{2+}], increase caused by Mch (Figure 3B). At the tissue level, FPL prevented chloroquine from relaxing Mch precontracted mouse ASM in a dose-dependent manner (Figure 3C). These results suggest that bitter tastants inhibit L-type \text{VDCCs, which in turn leads to a decrease in \([\text{Ca}^{2+}], and resulting bronchodilation.

**Bitter Tastants Reverse the \([\text{Ca}^{2+}], Rise and Contraction Evoked by Depolarization-Induced Activation of L-Type \text{VDCCs**

To directly examine the inhibitory role of bitter tastants on L-type \text{VDCCs, we studied the effect of bitter tastants on L-type \text{VDCC currents using patch clamp recording. Figure 4A shows two representative traces of L-type \text{Ca}^{2+} \text{ currents evoked by a depolarizing pulse to 0 \text{mV from a holding potential of } \pm 70 \text{mV (left panel) and mean peak currents at different depolarizing voltages (right panel) before and after 1 \text{mM} chloroquine. As is evident, chloroquine inhibited the L-type \text{Ca}^{2+} \text{ current when depolarizing voltages are between } \pm 30 \text{mV and } +40 \text{mV. To study whether this inhibition by chloroquine of L-type \text{VDCCs could produce relaxation, we evaluated the effect of this bitter tastant on the depolarization-induced increase in \([\text{Ca}^{2+}], and contraction. KCl is a standard and common reagent used to study cellular processes mediated by depolarization. In airways, depolarization is expected to not only activate \text{VDCCs in ASM but also those in cholinergic nerves as well [leading to release of Ach]. Indeed, in ferret and pig, KCl activates both mechanisms to cause airway contraction [40,41]. However, the Ach release mechanism does not operate in dog and rabbit airway, as demonstrated by several reports that showed that atropine, a muscarinic receptor antagonist, had no measureable effect on the magnitude of tension generated by high K\text{+ in these species [42–44]. Therefore to determine which mechanisms are activated by KCl in mouse airways, we examined the influence of atropine on KCl-induced contraction. Atropine dose-dependently inhibited Mch-induced contraction of mouse airways, and at concentrations greater than 100 \text{nM} it fully blocked the contraction (Figure S5A). These results confirm the efficacy of atropine in inhibiting muscarinic receptors in mouse airways. We further found that atropine (100 \text{nM}) reduced KCl (60 \text{mM})-induced contraction to 58\%±2.5\% of the time matched control (Figure S5), implying that KCl does activate \text{VDCCs in both ASM and cholinergic neurons to cause airway contractions via a combined effect. Although L-type \text{VDCCs are the major \text{Ca}^{2+} \text{ channel for \text{Ca}^{2+} \text{ influx upon depolarization in ASM, and mouse ASM cells exhibit only L-type \text{Ca}^{2+} \text{ currents [17], it is possible that KCl-induced contraction might involve Rho and Rho kinase via a \text{Ca}^{2+}-independent mechanism [45]. To examine this possibility, we studied the effect of extracellular \text{Ca}^{2+} \text{ on the KCl-induced increase in \([\text{Ca}^{2+}], and contraction. In \text{Ca}^{2+}-containing medium, KCl (60 \text{mM}) induced a marked increase in \([\text{Ca}^{2+}], (Figure S6A) in isolated ASM cells and airway contraction (Figure S6B). Yet in \text{Ca}^{2+}-free medium, the same KCl failed to cause any increase in \([\text{Ca}^{2+}], or a significant contraction (Figure S6B and S6C), consistent with published results in mouse and rat ASM [46,47]. These results indicate that KCl depolarizes the membrane, leading to a rise in \([\text{Ca}^{2+}], and a resultant contraction in mouse airway. They also demonstrate that \text{Ca}^{2+} \text{ influx is necessary to produce the KCl-induced contraction, and that a \text{Ca}^{2+}-independent mechanism (such as the suggested Rho and Rho kinase pathway [45,48,49] is not sufficient (if needed at all) to produce contraction. To further establish the role of L-type \text{Ca}^{2+} \text{ channels in the KCl-induced rise in \([\text{Ca}^{2+}], and contraction, we investigated the influence of diltiazem on these two effects of KCl. We found that 100 \mu\text{M} diltiazem pretreatment reduced the KCl-induced increase in AF/F0 from 122\%±19\% to 16.8\%±10\% in isolated ASM cells (n=9; Figure S6C), it also reversed the KCl-induced contraction by 93.1\%±4.8\% in airway tissue (n=6; Figure S6D). Therefore, in mouse ASM, high KCl seems to increase \([\text{Ca}^{2+}], and cause contraction by depolarizing the membrane and activating L-type \text{VDCCs. Considering the action of KCl as just described (Figures S3 and S6), we reasoned that bitter tastants would be able to relax airways precontracted by KCl if bitter tastant’s inhibition of L-type \text{Ca}^{2+} \text{ channels underlies its relaxation of airways pre-contracted by Mch (Figure 3). Indeed, we found that a 60 \text{mM} KCl-induced contraction in mouse and human airways was fully reversed by...
either chloroquine (1 mM) or denatonium (1 mM) (Figures 4B and S2C). This reversal is due, at least in part, to a direct inhibition of VDCCs in ASM because (1) in the presence of 100 nM atropine, chloroquine can fully block atropine-resistant contraction (Figure 4B), and (2) when nerve action potentials were blocked by 1 μM tetrodotoxin, a voltage-dependent Na+ channel blocker, and arachidonic acid metabolism was inhibited by 1 μM indomethacin, a nonselective inhibitor of cyclooxygenase, the bitter tastants still fully relaxed airways precontracted by KCl (Figure 4B). Similar to their effects on Mch-induced responses (Figure 2B), chloroquine reversed the KCl-induced increase in [Ca2+]i, and shortening of isolated single ASM cells (Figure 4C; n = 7). Moreover, FPL dose-dependently reversed chloroquine-induced relaxation in ASM pre-contracted by KCl (60 mM; Figure 4D), and prevented the reduction of [Ca2+]i, by chloroquine in cells stimulated by KCl (Figure 4E).

Gβγ Activation Mediates Bitter Tastant Suppression of the Rise in [Ca2+]i, Evoked by Activation of L-Type VDCCs

To address the signaling basis underlying bitter tastant inhibition of L-type VDCCs, we studied the impact of perturbing TAS2R signaling on bitter tastant-induced reversal of the [Ca2+]i increase in response to KCl in isolated single ASM cells. Pretreatment with PTX at 1 μg/ml for 6-8 h prevented chloroquine-induced reversal of the KCl-induced increase in [Ca2+]i, as did gallin (20 μM) and anti-βγ, a Gβγ blocking peptide (1 μM) (Figure 5). However, U73122 and 2-ABP, at the concentrations that block the bitter tastant-induced increase in [Ca2+]i, in resting cells (Figure 1), failed to alter chloroquine’s ability to reverse a KCl-induced increase in [Ca2+]i (Figure 5). These results indicate that activation of Gβγ but not PLCβ and IP3R is required for bitter tastant-induced inhibition of L-type VDCCs.

Discussion

Our results demonstrate that bitter tastant’s reversal of the rise in [Ca2+]i, evoked by bronchoconstrictors is required for its bronchodilation effect. They also reveal that bitter tastants can generate different and opposing Ca2+ signals depending upon the cellular environment. When administered alone to ASM cells at rest, bitter tastants activate the canonical TAS2R signaling pathway to modestly raise [Ca2+]i (Figure 5C) without affecting the contraction. Yet when applied in the presence of the bronchoconstrictors Mch and KCl, they inhibit L-type VDCCs, leading to a reversal of both the evoked [Ca2+]i rise and the contraction (Figure 5C). Remarkably, both types of Ca2+ signals require Gβγ, while only the increase in resting [Ca2+]i, depends on PLCβ2 activation and IP3 generation.

Bitter taste receptors (35 in mouse and 25 in human) belong to seven transmembrane domain G-protein-coupled receptors. Long thought to only be expressed in the epithelium cells of the taste buds of the tongue, recent studies have revealed that these receptors also express in several extraoral tissues including brain, testis, immune cells, gastrointestinal tract, and respiratory system [50–57]. In airways, these receptors are found to be expressed in ciliated epithelial cells and nasal solitary chemosensory cells [51,52]. Deshpande et al. [6] reported that multiple TAS2Rs can be detected in cultured human ASM cell lines. In this study, for the first time, to our knowledge, we found that this class of G-protein-coupled receptors is expressed in native mouse ASM cells. Specifically, we determined that TAS2R107, to which both chloroquine and denatonium are ligands, localizes in the cell periphery, a location well suited for mediating bronchodilation in response to these two bitter tastants. We cannot rule out that other types of TAS2Rs also contribute to the bronchodilation induced by chloroquine and denatonium, since both of them can activate multiple mouse and human TAS2Rs [58]. Nevertheless, these ligands do activate TAS2R signaling transduction, resulting in a bronchodilation effect, because pharmacological blocking of multiple downstream components of bitter taste receptors can prevent chloroquine and denatonium-induced cellular responses (Figures 1C, 5A, and 5B). It is also worth noting that chloroquine, denatonium, and all the bitter tastants examined so far are not endogenous ligands for bitter taste receptors. Hence a major question remains as to whether bitter taste receptors in ASM cells have physiological ligands. Interestingly, a recent study revealed that acyl-homoserine lactones, quorum-sensing molecules for Gram-negative pathogenic bacteria, can activate bitter taste receptors in nasal solitary chemosensory cells to evoke trigeminally mediated reflex reactions, which may trigger an epithelial inflammatory response before the bacteria reach population densities capable of forming destructive biofilms [52,53]. It would be of great interest and significance to investigate whether these quorum-sensing molecules can activate bitter taste receptors in ASM to induce bronchodilation.

This study revealed two major differences in Ca2+ signaling compared to the study by Deshpande et al. [6]. First, these authors reported that bitter tastant increased [Ca2+]i, to a level comparable to bronchoconstrictors. In freshly isolated ASM, we found that bitter tastants only modestly increase [Ca2+]i, to a level much lower than that produced by bronchoconstrictors. Second, Deshpande et al. [6] reported that bitter tastants generate local Ca2+ events. However, in freshly isolated ASM, we found that bitter tastants do not increase local Ca2+ releases such as Ca2+ puffs and Ca2+ sparks. A reason for these two discrepancies may be that Deshpande et al.’s studies were conducted in cultured ASM cell lines; compared to freshly isolated ASM, these cells display a different phenotype by altering the expression of receptors, ion channels, and contractile proteins [13,14]. The aforementioned two differences and another difference in which we found that bitter tastants do not activate large-conductance Ca2+-activated K+ channels strongly argue that bitter tastant-induced bronchodilation is highly unlikely to result from the generation of local Ca2+ events, which in turn activate large-conductance Ca2+-activated K+ channel and hyperpolarize the membrane as proposed [6].

Since bitter tastants relax precontracted airways [6,11,15,16], it is imperative to use a similar stimulating paradigm in order to understand the underlying mechanism of this relaxation. By simultaneously measuring [Ca2+]i, and cell shortening, we found that bitter tastant’s ability to reverse the increase in [Ca2+]i, caused by bronchoconstrictors is the underlying signal producing the bronchodilation. Three lines of evidence support this conclusion. First, in the presence of bronchoconstrictors, bitter tastants lowered [Ca2+]i, while at the same time relaxing the precontracted cells, and this response was reversible. Second, clamping intracellular [Ca2+]i, to levels produced by the bronchoconstrictors (low μM) prevented bitter tastants from relaxing airways. Third, enhancing and blocking Ca2+ influx via L-type Ca2+ channels can oppositely regulate the relaxation mediated by bitter tastants. These results reinforce the idea that [Ca2+]i, is the critical signal governing ASM contractility.

The opposing Ca2+ signals mediated by Gβγ upon activation of TAS2Rs revealed in this study are unique. It is expected that gustducin Gβγ activates PLCβ to generate IP3 and release Ca2+ from endo/sarcoplasmic reticulum to raise [Ca2+]i, in ASM cells. But, unexpectedly, gustducin Gβγ also suppresses Ca2+ signaling mediated by Mch, which largely activates M5R, a Gq family.
βγ-Gustducin Inhibits L-Type Ca^{2+} Channel

A

![Graph showing current (I_{Ba}) in pA at different voltages (0 mV).](image)

B

![Graph showing tension in mN vs. time (10 min).](image)

C

![Graph showing F/F₀ (%) vs. time (20 s).](image)

D

![Graph showing tension in mN vs. log [FPL] (M).](image)

E

![Graph showing normalized ΔF/F₀ vs. time (10 s).](image)
Figure 4. Bitter tastants block L-type VDCCs. (A) Chloro blocked L-type VDCC currents. Left panel displays patch clamp recordings of L-type Ca\(^{2+}\) currents in response to a voltage pulse from \(-70\) mV to \(0\) mV in the control and in the presence of \(1\) mM chloro, and the right panel the effect of chloro on the current-voltage \(I-V\) relationship of the Ca\(^{2+}\) current \((n=5)\). Ba\(^{2+}\) was used as a charge carrier, and the peak current was used to construct the \(I-V\) relationship. Note that the high peak potential for activation seen in the \(I-V\) relationship, and its sensitivity to FPL and nifedipine [17] indicate these Ca\(^{2+}\) currents resulted from the opening of L-type VDCCs. (B) Bitter tastants relaxed KC-induced contraction of mouse airways. The left panel shows representative force recordings in response to 60 mM KCl followed by \(1\) mM chloro in the presence of \(100\) mM atropine and in its absence. The right panel shows the mean values of the relaxation of KC-induced contraction by \(1\) mM chloro in the control \((n=9)\), in the presence of \(100\) mM atropine \((n=16)\), or in the presence of \(1\) mM tetrodotoxin (TTX) and \(1\) mM indomethacin (Indom) \((n=6)\). Note that the inhibition of chloro in the presence of \(100\) mM atropine was calculated relative to the atropine-resistant contraction in response to KC. Not shown: \(1\) mM denatonium relaxed airways precontracted by \(60\) mM KCl by 97\% \(\pm\) 4\% \((n=5\) independent experiments). (C) Relationship between \([\text{Ca}^{2+}]_i\) and cell length in response to \(60\) mM KCl and \(1\) mM chloro. Left panel shows the time course of concomitant changes in \([\text{Ca}^{2+}]_i\) and cell length and the right the mean \(\pm\) SEM \((n=15)\) at the four time points marked on the left. \(p<0.05; \ast p<0.01\). (D) FPL 64176 (FPL) dose-dependently inhibited chloro-induced bronchodilation of KC precontracted airways. Left panel shows two representative recordings, and the right panel the means \(\pm\) SEM \((n=5–7\) independent experiments). \(60\) mM KCl and \(1\) mM chloro were used. Given the non-monotonic nature of the relaxation (left), both the greatest reduction in force after chloro \((\text{i.e., maximum})\) and the force reduction \(5\) min after chloro were measured and divided by the peak force: resting force before application of chloro. (E) FPL inhibited chloro-induced suppression of the rise in \([\text{Ca}^{2+}]_i\), produced by KCl. Left panel shows original recordings of Ca\(^{2+}\) responses and the right panel the mean \(\pm\) SEM \((n=28\) without FPL, \(n=16\) with FPL). The values represent as \((\Delta F/F_0)\) at the peak after KC – AF/F\(_0\) at 30 s after chloro)/(AF/F\(_0\) at the peak after KC – \(\Delta F/F_0\) at basal) \(>\times\)100.

doi:10.1371/journal.pbio.1001501.g004

**Beta-gustducin Inhibits L-Type Ca\(^{2+}\) Channel**

In general, \(\beta\gamma\) from the \(G_\gamma/G_\alpha\) family (to which TAS2Rs belong) tends to potentiate, rather than, inhibit the Ca\(^{2+}\) responses caused by the \(G_\gamma\) family [59,60]. It remains to be determined whether the inhibition of Ca\(^{2+}\) signaling by TAS2R activation is \(\beta\gamma\) isoform specific. Since \(\beta\gamma\) also mediates activation of M2R and \(\gamma\)-aminobutyric acid-B receptors [61,62], our present findings suggested that \(\beta\gamma\) reversal of the rise in \([\text{Ca}^{2+}]_i\), caused by bronchonstrictors is isoform specific, and is likely via \(\beta\gamma\beta\gamma\gamma\) dimers, which are released upon activation of TAS2Rs [63]. Further studies using ASM cells with genetic deletions of these isoforms should facilitate studying this possibility. It is worthy of mention that virtually all of the studies of bitter taste signaling in taste buds [7–10] and extraoral tissues [51–53] have focused on the responses mediated by bitter tastants alone; the opposing Ca\(^{2+}\) activation is \(\beta\gamma\) isoform specific. Since \(\beta\gamma\) is likely that \(\beta\gamma\) also mediates the effect of bitter tastants and other activators.

L-type VDCCs in smooth muscle can be modulated by a variety of means including phosphorylation and Ca\(^{2+}\) [64–68]. Yet for the first time, to the best of our knowledge, we show that \(\beta\gamma\) subunits of G-protein gustducin can inhibit these channels in smooth muscle, extending the similar findings for cloned Cav 1.2 in heterologous expression cells and Cav1.1 in skeletal muscle fiber [69,70]. This interpretation is buttressed by the experiments showing that the contraction mediated by KC-induced activation of presynaptic Ca\(^{2+}\) channels can also be fully blocked by bitter tastants (Figure 4B). What remains unknown is whether \(\beta\gamma\) directly or indirectly inhibits these channels, and the structural basis for this inhibition. Given that \(\beta\gamma\) can directly inhibit \(K^+\) channels and N-type Ca\(^{2+}\) channels in several cell types [71–75], it is likely that \(\beta\gamma\) acts on L-type VDCCs in a similar manner.

Gustducin \(\beta\gamma\) subunits inhibit L-type VDCCs to cause bronchodilation, highlighting the importance of these channels in mediating bronchoconstriction and their potential as a target for bronchodilators. Indeed, L-type VDCCs are expressed in ASM cells and their activation causes these cells to fully contract (Figures 4, S4, S5, S6) [17,27,76,77]. Also, activation of these channels is a major mechanism underlying bronchoconstrictor-induced contraction of different species including airway and human [25,27,30,32], but see [24]). Moreover, three classes of organic L-type VDCC blockers (i.e., dihydropyridines, phenylalkylamines, and benzothiazepines) are effective in relieving airway spasm in animal models of asthma and in exercise-induced asthmatic patients [78–81]. A long-standing puzzle regarding L-type channels in ASM is that clinical trials in the 1980s suggested that antagonists for this channel were of limited use treating asthma in the population as a whole [81,82]. A potential reason for this enigma may be to some extent related to the mode of action of these blockers. It is known that these classic organic blockers exert their inactivation of L-type VDCCs in a voltage, stimulation, and frequency dependent manner [34,36,37]. Interestingly, allergen sensitized guinea-pig and rabbit ASM cells have a more hyperpolarized membrane potential than normal cells [83]. This implies that L-type Ca\(^{2+}\) channel blockers (for example, dihydropyridines) would bind more weakly with these channels, thus decreasing the efficacy of these agents to inhibit these channels, should ASM cells from asthma patients have a more negative membrane potential. Bitter tastants, by their ability to inhibit L-type Ca\(^{2+}\) channels via activation of gustducin \(\beta\gamma\), perhaps could circumvent the drawbacks of the currently available L-type Ca\(^{2+}\) channel blockers, and thus be a more effective asthma treatment. This is likely given that bitter tastants induce a stronger bronchodilation in both in vitro and in vivo asthmatic mouse models than do \(\beta\) agonists [6,11], the most commonly used bronchodilators for treating asthma and COPD.

Although bitter tastants are promising candidates to be developed as a new class of bronchodilators, and the findings in the present study provide the cellular and molecular rationale for this line of inquiry, we would caution that chloroquine and denatonium examined in this study may not be ideal candidates because of the high concentration (i.e., on the order of 100 \(\mu\)M) needed to fully relax precontracted ASM. This caveat, however, should not dampen enthusiasm for this endeavor as there are many thousands of bitter tastants available from plants and animals, and numerous bitter small molecules synthesized by research laboratories and a variety of companies over the years. In fact, bitter tastants can stimulate bitter taste receptors at concentrations in the nanomolar range: strychnine activates human TAS2R46 with an EC\(_{50}\) of 430 nM and aristolochic acid activates human TAS2R43 with an EC\(_{50}\) of 81 nM [84,85]. Therefore, it is highly likely that bitter tastants with a highly potent bronchodilating action can be discovered. Searching for these bitter tastants is of clinical significance because the current bronchodilators are insufficient for treating severe asthma and many COPD patients. A critical step in identifying highly potent bitter tastants is developing reliable and highly effective screening methodologies. Simultaneous measurements of cell shortening and the \([\text{Ca}^{2+}]_i\) signal (i.e., a decrease of elevated \([\text{Ca}^{2+}]_i\)) as developed in the present study, are robust and quantitative and provide a powerful paradigm for identifying potential bronchodilators from among the many bitter tastants available.
Materials and Methods

Animal Tissue Handling

Experimental protocols for animal research were approved by the Institutional Animal Care and Use Committees at the University of Massachusetts Medical School (protocol A-1473 to RZG).

Isolation of Mouse Airway Smooth Muscle Cells

C57BL/6 mice from 7 to 12 wk of age were anesthetized with intraperitoneally injected pentobarbitone (50 mg kg\(^{-1}\)), and the trachea and mainstem bronchi were quickly removed and placed in a pre-chilled dissociation solution consisting of (in mM): 135 NaCl, 6 KCl, 5 MgCl\(_2\), 0.1 CaCl\(_2\), 0.2 EDTA, 10 HEPES, and 10 Glucose (pH 7.3). Tracheas and mainstem bronchi were dissected free from the surface of the connective tissue. The airway tissue was incubated in the dissociation medium containing papain 30 unit/ml, 1 mM DTT, and 0.5 mg/ml BSA, at 35°C for 30 min, and then transferred to a dissociation medium containing 3 unit/ml collagenase F and 0.5 mg/ml BSA, and incubated at 35°C for another 15 min to produce isolated ASM cells. Finally, the tissue was agitated with a fire polished wide-bore glass pipette to release the cells.

Mouse Airway Smooth Muscle Contraction Bioassay

C57BL/6 mice at 7–12 wk of age were sacrificed and the entire respiratory trees were rapidly removed and immersed in Krebs physiologic solution containing (in mM) 118.07 NaCl, 4.69 KCl, 2.52 CaCl\(_2\), 1.16 MgSO\(_4\), 1.01 NaH\(_2\)PO\(_4\), 25 NaHCO\(_3\), and 11.10 glucose. Trachea and mainstem bronchi were isolated and cut into rings (4 mm in length). The rings were mounted on a wire myograph chamber (Danish Myo Technology), and a PowerLab recording device (AD Instruments) was used to record isometric tension. The ring preparations with zero tension were immersed in 5 ml of Krebs physiologic solution, bubbled with 95% O\(_2\) and 5% CO\(_2\) at 37°C. After 10 min equilibration, three stretches (each 2.5 mN) at 5 min intervals were applied to the rings. After these stretches the basal tones of the rings were usually settled at approximately 2 mN. To test the contractile response, each ring was stimulated twice with KCl (60 mM), separated by 20 min, before proceeding to other treatments. The order and treatment time of agonists and antagonists are indicated in the figure captions. In the experiments in Figures 4B, S4B, and S4D, 1 mM tetrodotoxin was added to prevent action potentials of neurons and 1 mM indomethacin to inhibit cyclooxygenase. The force in response to 60 mM KCl in the presence of tetrodotoxin and indomethacin was 95% ± 7% (n = 6) of that in their absence.

Airway Smooth Muscle Permeabilization

Mouse bronchi rings (4 mm in length) free of connective tissues were incubated for 5 min in HEPES-Tyrode (H-T) buffer which contained 137.0 mM NaCl, 2.7 mM KCl, 1.0 mM MgCl\(_2\), 1.8 mM CaCl\(_2\), 10 mM HEPES, 5.6 mM glucose (pH 7.4). The rings were then transferred to and incubated in Ca\(^{2+}\)-free H-T buffer for 5 min followed by another 5 min in buffer A (30 mM TES, 0.5 mM DTT, 50 mM KCl, 5 mM K\(_2\)EGTA, 150 mM

Figure 5. Bitter tastants inhibit L-type VDCCs via a G\(_{\beta\gamma}\) dependent process. (A) Representative recordings of changes in [Ca\(^{2+}\)] in response to 60 mM KCl followed by chloro (1 mM) with and without pretreatment with PTX (1 µg/ml), gallein (1 µM), anti-βγ blocking peptide (1 µM), U73122 (3 µM), and 2-APB (50 µM). The application protocols for these compounds were the same as in the experiments in Figure 1C. All data were scaled to have a maximum of 100 and aligned at the time point when KCl was administrated. (B) Effects of compounds listed in A on chloro-induced suppression of KCl-induced increase in [Ca\(^{2+}\)]. The values were calculated the same as in Figure 4E. Compared to the control (i.e., chloro alone after KCl, black filled bar), p<0.0001 for PTX, gallein, and anti-βγ; and p>0.05 for U73122 and 2-APB. Data are shown as mean ± SEM (n = 12–38 cells). (C) A model for TAS2R signaling and bitter tant-induced bronchodilatation. We propose that bitter tastants activate the canonical TAS2R signaling cascade; this modestly increases [Ca\(^{2+}\)] in resting cells but exerts no significant effect on resting tone. On the other hand, activation of TAS2Rs activates gudstducin and release G\(_{\beta\gamma}\), which turns off L-type VDCCs that are pre-activated by bronchoconstrictors, leading to bronchodilatation.

doi:10.1371/journal.pbio.1001501.g005

Materials and Methods
sucrose [pH 7.4]). To skin ASM, the rings were incubated for 45 min with α-toxin (16,000 units/ml) in buffer A at room temperature. After the permeabilization, the rings were treated with 10 μM ionomycin for 10 min to deplete intracellular Ca^{2+} stores.

Skinned airway rings were mounted on the wire myography chamber and washed two times with pCa 9 solution (20 mM TES, 4 mM K2EGTA, 5.83 mM MgCl2, 7.56 mM potassium propionate, 3.9 mM Na2ATP, 0.5 mM dithioerythritol, 16.2 mM phosphate creatinine, 15 units/ml creatine kinase [pH 6.9]). The viability of the skinned muscle rings was examined by stimulation with pCa 4.5 solution (20 mM TES, 4 mM CaEGTA, 5.66 mM MgCl2, 7.53 mM potassium propionate, 3.9 mM Na2ATP, 0.5 mM dithioerythritol, 16.2 mM phosphate creatinine, 15 units/ml creatine kinase [pH 6.9]) followed by the pCa 9.0 solution. The muscles that could generate sustained contraction in response to the pCa 4.5 solution and fully relax when exposed to the pCa 9.0 solution were used for subsequent experiments. To test the effect of bitter tastants, the viable muscle rings were induced to contract by exposure to pCa 5.5 solution followed by the administration of bitter tastants at the concentrations indicated in Figure 3.

**Measurement of Global [Ca^{2+}] and Ca^{2+} Sparks**

Fluorescence images using fluo-3 as a calcium indicator were obtained using a custom-built wide-field digital imaging system. The camera was interfaced to a custom made inverted microscope, and the cells were imaged using either a 20× Nikon 1.3 NA for global [Ca^{2+}] measurement or a 60× Nikon 1.4 NA oil for Ca^{2+} spark measurement. The 488 nm line of an Argon Ion laser provided fluorescence excitation, with a shutter to control exposure duration, and emission of the Ca^{2+} indicator was monitored at wavelengths >500 nm. The images were acquired at the speed of either 1 Hz for global [Ca^{2+}] measurement or 50 Hz for Ca^{2+} spark measurement. Subsequent image processing and analysis was performed off line using a custom-designed software package, running on a Linux/PC workstation. [Ca^{2+}] was represented as ΔF/F_0×100 with F calculated by integrating fluo-3 over entire cells for global [Ca^{2+}] after background correction with areas free of cells, or just the value at the brightest pixel (i.e., epicenter pixel) for Ca^{2+} sparks.

**Patch-Clamp Recording**

Membrane currents were recorded with an EPC10 HEKA amplifier under perforated whole-cell patch recording configuration. The extracellular solution contained (in mM): NaCl 126, tetraethylammonium Cl 10, BaCl2 2.2, MgCl2 1, Hepes 10, and glucose 5.6 [pH adjusted to 7.4 with NaOH]. The pipette solution contained (in mM): CaCl2 139, MgCl2 1, Hepes 10, MgATP 3, Na2ATP 0.5 [pH adjusted to 7.3 with KOH]; amphotericin B was freshly made and added to the pipette solution at a final concentration of 200 μg/ml. Whole-cell Ba^{2+} currents were evoked by step depolarization with 300 ms duration every 10 s from a holding potential of −70 mV at a 10 mV increment (Figure 4A) or with protocol as described in the caption of figure caption (Figure S4C). Currents were leak corrected using a P/4 protocol.

**Measurement of Cell Shortening**

Myocytes were placed into a recording chamber superfused with the bath solution for patch clamp experiments at room temperature. Cells loaded with Fluo-3 were imaged using a custom-built wide-field digital imaging system and their lengths were determined using custom software to manually trace down the center of the cell [17].

**Reverse Transcription-PCR to Detect mRNA**

The connective tissues in trachea and mainstem bronchi were carefully removed and the ASM were then quickly frozen in dry ice. The total RNA of the ASM was isolated with the TRIzol (Invitrogen) method following the manufacturer’s guidelines; cDNA was synthesized using extracted RNA with an Omniscript Reverse Transcription kit (Qagen). The specific primers, synthesized by Invitrogen, are listed in Table S1. β-actin was used as a positive control and the absence of DNA as a negative control, and the PCR reactions were carried out in a PCR mastercycler.

**Immunocytochemistry**

Mouse ASM cells, isolated as described above and plated onto poly-L-lysine coated coverslips were fixed and permeabilized (0.1 M ethanolamine in PBS plus 0.1% triton X-100 [pH 8]) and then immunolabeled as described previously [86]. Anti-TAS2R107, an affinity purified rabbit polyclonal antibody raised against a peptide mapping within an extracellular domain of mouse TAS2R107, was purchased from Santa Cruz Biotechnol (sc-139175), and purified IgG was used as control.

**3D fluorescence imaging was performed on an inverted wide field microscope (Nikon Diaphot 200) with excitation by a 100 W mercury lamp. Images were obtained through a 60× objective and digitally recorded on a cooled, back-thinned CCD camera (Photometrics), with an effective pixel size at the specimen of 83 nm in x-y and a z spacing of 100 nm. This resulted in a 3D stack of approximately 100 image planes for each cell. The fluorescence images were deconvoluted with a constrained, iterative approach [87] originally designed for UNIX systems. The algorithm was rewritten using FFTW, a free, fast Fourier transform library and implemented as a multiuser client/server system on computers running the Fedora operating system (Red Hat), either stand-alone or configured in a Beowulf cluster. Each image was dark current and background subtracted, flat-field corrected, and then deconvolved. After deconvolution images were thresholded to eliminate non-specific binding. Voxels that fell below a threshold were considered to be non-specific bindings and were set to zero; all other voxels remained unchanged. This threshold was derived from analysis of control images containing purified IgG. The intensity which eliminated 99% of the voxels in the control images became the threshold intensity.

**Reagents and Their Application**

All chemicals, except fluo-3 (Invitrogen Co), gallein (Tocris Bioscience), anti-β2 blocking peptide (AnaSpec), anti-TAS2R107 (Santa Cruz Biotechnology), and purified IgG (Jackson Immunoresearch Laboratories) were purchased from Sigma-Aldrich Co. For single cell studies, agonists and antagonists were applied locally to cells via a picospritzer at a constant pressure, so that the duration of its action and concentration could be controlled easily.

**Statistics**

Unless stated otherwise, data are reported as mean ± standard error of the mean (SEM) and n represents the number of cells or trachea and mainstem bronchi. Statistical analysis of differences was made with Student’s paired or unpaired t-test and the significance level was set at p<0.05.

**Supporting Information**

Figure S1 Bitter tastant chloroquine dose-dependently increased [Ca^{2+}], in resting single cells (A) without a significant effect on the contractility (B) of relaxed mouse airways. Results are mean ± SEM, (n = 5–30 cells in...
Figure S2 Characteristics of [Ca^{2+}]_i and contractile responses to bitter tastants and diltiazem in human ASM. (A) Bitter tastants reversed the [Ca^{2+}]_i rise and cell shortening induced by Mch. Measurements were taken at the steady state levels in response to Mch and chloroquine. The cell length before stimulation was considered as 100%. *: Student's t-test; ***: p<0.001; n = 6–12. (B) L-type VDCC blocker diltiazem dose-dependently reversed 10 μM Mch-induced contraction (n = 5 independent experiments). % relaxation = tension decrease due to diltiazem divided by tension increase due to Mch, times 100. The tension decrease at each concentration of diltiazem is measured once the tension stabilizes. The tension decrease at each increased concentration is always measured relative to the peak tension (i.e., it is total decrease, not the incremental decrease due to the additional diltiazem which was added). (C) Chloroquine (1 mM) and diltiazem (100 μM) relaxed human intrapulmonary bronchi precontracted by 60 mM KCl (n = 3–5 independent experiments). % relaxation = tension decrease due to chloroquine divided by tension increase due to Mch, times 100. Bar charts are mean ± SEM. Human lung tissues were obtained (with informed consent) from patients undergoing surgery (lobectomy) for lung cancer at the Department of Surgery and the Department of Pathology at the University of Massachusetts Memorial Medical Center (Worcester). The tumors were identified as non-small cell carcinoma (adenocarcinoma or squamous cell carcinoma). Intrapulmonary airways were dissected out and cleaned free of the connective tissues. These airways were either cut into the rings (4 mM in length) for force measurements the same as for mouse airway rings, or digested with the same enzymes, dissociation medium, and isolation procedures as for single mouse ASM cells. The experimental protocols on human tissues were approved by the Committee for Protection of Human Subjects in Research at the University of Massachusetts Medical School (protocol 13590 to RZG). (TIFF)

Figure S3 Ca^{2+} influx plays a major role in producing and maintaining Mch-induced increases in [Ca^{2+}]_i and contraction in mouse ASM. (A) In Ca^{2+} free medium, the tension generated by Mch was less than 20% of that in the presence of extracellular Ca^{2+}. ***: p<0.001, Student’s paired t-test, n = 9 airway rings for the group with Ca^{2+}, and n = 10 airway rings for the group without Ca^{2+}. (B) Mch increased [Ca^{2+}], much less in Ca^{2+} free medium than in the presence of extracellular Ca^{2+}. In the absence of extracellular Ca^{2+}, Mch (10 μM) produced different patterns of changes in [Ca^{2+}]_i, so the area under each curve was calculated for 1 min of Mch stimulation and compared between the two conditions (right panel). **: p<0.001 with extracellular Ca^{2+} (n = 9 cells) versus without the Ca^{2+} (n = 12 cells); Student’s unpaired t-test. (C) Ca^{2+} stores remained functional in the absence of extracellular Ca^{2+}. The cells were placed in the absence of extracellular Ca^{2+} for ~15 min, and then stimulated with two 10 μM Mch pulses 15 min apart. The chart on the right indicates that two Mch administrations produced comparable Ca^{2+} responses, i.e., Ca^{2+} stores are intact under experimental conditions in the present study. NS, p>0.05 for the response in the first pulse of Mch versus that in the second pulse, Student’s paired t-test, n = 10. ΔF/F₀ for (B) and (C) are the average over the entire cell. (TIFF)

Figure S4 Role of L-type Ca^{2+} channel activation in Mch-induced contraction in mouse airway. (A) Left panel: L-type VDCC blocker diltiazem dose-dependently reversed Mch-induced contraction (using tension as a proxy measure). Right panel: results for n = 6 airway rings. % relaxation was calculated the same as in Figure S2B. (B) Diltiazem inhibited Mch- and KCl-induced contraction in a dose-dependent manner. (i) Once equilibrated, airway rings generated stable responses to 60 mM KCl (i.e., K₀, K₁, and K₂), and to 1 μM Mch (i.e., M₀ and M₁) over a time span longer than 1 hr. (ii, iii, iv) show representative responses to KCl and Mch in the presence of diltiazem at 1 μM, 10 μM, and 100 μM, respectively. Two pulses of KCl (K₁ and K₂) were administrated before Mch (M₁) to facilitate diltiazem inactivation of L-type Ca^{2+} channels. (v) Dose-response curve for diltiazem-mediated inhibition of contraction by KCl. Data are mean ± SEM (n = 5); % inhibition = (force by M₀ – force by K₂)/(force by K₀)×100. (vi) Dose-response curve for diltiazem-mediated inhibition of contraction by Mch. Data are mean ± SEM (n = 5); % inhibition = (force by M₀ – force by M₁)/(force by M₀)×100. Forces are measured at their peak, excluding the noise spikes due to the washing out of KCl or Mch. (C) Diltiazem inhibited L-type VDCC currents in a dose-dependent manner. Diltiazem was added to the bath solution cumulatively. Once at equilibrium at each level of diltiazem, cells were stimulated with a train of ten voltage pulses from ~70 mV to 0 mV (inset) at 10 s intervals. Left panel displays patch clamp recordings of L-type Ca^{2+} currents in the control and in the presence of diltiazem at the given concentration (each in response to the tenth voltage pulse), and the right panel depicts the effect of diltiazem on the peak current at different concentrations. Ba^{2+} was used as the charge carrier. Data are mean ± SEM (n = 6); % inhibition = peak current of the control – peak current at given diltiazem concentration)/peak current of the control ×100 (measured in the tenth voltage pulse). (D) Effect of nisoldipine on Mch-induced contraction. In light of the voltage-dependence of the inhibition of nisoldipine on L-type VDCCs, airway rings were modestly depolarized with 20 mM KCl (red bars) before Mch stimulation. Left panel displays a representative contractile response to 60 mM KCl and 1 μM Mch before and after 1 μM nisoldipine. Right panel shows the mean values (mean ± SEM) of inhibition by 1 μM nisoldipine of contraction evoked by KCl and Mch, respectively. % inhibition = (maximal force at the control – maximal force with nisoldipine)/maximal force at the control ×100. (TIFF)

Figure S5 KCl activates VDCCs in cholinergic nerves and ASM. (A) Atropine dose-dependently inhibited Mch-induced mouse airway contraction. The left panels display representative contractile responses to 3 μM Mch in the absence or the presence of atropine as marked near the traces, and the right panel shows the mean values (mean ± SEM; n = 4–8) of inhibition by atropine of contraction evoked by 3 μM Mch. % inhibition = (maximal force from the control – maximal force with atropine)/maximal force from the control ×100. (B) Atropine inhibited KCl-induced mouse airway contraction. The left panel shows a representative tension recording in response to 60 mM KCl before and after 100 nM atropine, as marked beneath. The right panel shows the summarized results as the ratio of the force generated by the second KCl pulse over first KCl pulse (mean ± SEM; n = 6 for the time matched controls, n = 28 for atropine). (TIFF)

Figure S6 KCl activates L-type VDCCs to increase [Ca^{2+}]_i and cause contraction in mouse ASM. (A) KCl failed to generate any global [Ca^{2+}]_i increase in the absence of
extracellular Ca\(^{2+}\) in isolated single ASM cells. (i) A representative [Ca\(^{2+}\)]\(_i\) response to 60 mM KCl in the presence of extracellular Ca\(^{2+}\), (ii, iii, iv) three examples showing that the same concentration of KCl did not increase Ca\(^{2+}\) in the zero Ca\(^{2+}\) medium. This failure was not due to the depletion of intracellular Ca\(^{2+}\); it was because 10 \(\mu\)M Mch still induced Ca\(^{2+}\) release either as a single peak or as oscillations. Eight cells gave rise to similar responses. \(\Delta F/F_0\) is the average over the entire cell. (B) KCl (60 mM) caused virtually no increase in tension in the absence of extracellular Ca\(^{2+}\). The airways were placed in the Ca\(^{2+}\)-free solution for 15 min before the measurement commenced. Left panel shows a pair of representative recordings and right panel the average results. **p < 0.01, Student’s paired t-test, \(n = 6\) independent experiments. (C) KCl (60 mM)-induced increase in [Ca\(^{2+}\)]\(_i\), was markedly inhibited by prior application of L-type VDCC blocker diltiazem (100 \(\mu\)M). **p < 0.01, Student’s paired t-test, \(n = 9\) for each conditions. (D) Diltiazem (100 \(\mu\)M) relaxed 60 mM KCl-induced contraction of mouse airways. Data are mean ± SEM (\(n = 6\) independent experiments), and % relaxation definition and analysis are the same as in Figure S2C.

**Table S1 Primers for Reverse Transcription-PCR.**

**References**

1. Herschenson MB, Brown M, Cameretti-Mercado B, Sohayi J (2008) Airway smooth muscle in asthma. Annu Rev Pathol 3: 323–555.
2. Grainge CL, Lau LC, Ward JA, Dalay V, Lalith G, et al. (2011) Effect of bronchoconstriction on airway remodeling in asthma. N Engl J Med 364: 2006–2015.
3. Tibia O, Panettieri RA, Jr. (2009) Noncontractile functions of airway smooth muscle cells in asthma. Annu Rev Physiol 71: 509–535.
4. Fanta CH (2009) Asthma. N Engl J Med 360: 1002–1014.
5. Han MK, Martinez FJ (2011) Pharmacotherapeutic approaches to preventing acute exacerbations of chronic obstructive pulmonary disease. Pro Am Thorac Soc 8: 356–362.
6. Deshpande DA, Wang WC, McIlmoyle EL, Robinett KS, Schillinger RM, et al. (2010) Bitter taste receptors on airway smooth muscle broncholate by localized calcium signaling and reverse obstruction. Nat Med 16: 1299–1304.
7. Wong GT, Gannon KS, Margoluk RF (1996) Transduction of bitter and sweet taste by gustducin. Nature 381: 796–800.
8. Chandraharak J, Mueller KL, Hoon MA, Adler E, Feng L, et al. (2000) T2R function as bitter taste receptors. Cell 100: 703–711.
9. Zhang Y, Hoon MA, Chandraharak J, Mueller KL, Cook B, et al. (2003) Coding of sweet, bitter, and umami tastes: different receptor cells sharing similar signaling pathways. Cell 112: 293–301.
10. Kruz-Avila L, McLaughlin SK, Wildman D, McKinnon PJ, Robichon A, et al. (1995) Coupling of bitter receptor to phosphodiesterase through transducin in taste receptor cells. Nature 376: 80–85.
11. Zhang CH, Chen C, Lifshitz LM, Fogarty KE, Zhu MS, et al. (2012) Activation of BK channels may not be required for bitter-tastant-induced bronchodilation. Nat Med 18: 641–650.
12. Somoly AP, Somoly AV (1994) Signal transduction and regulation in smooth muscle. Somoly Physiol 72: 231–236.
13. Chandley-Campbell J, Campbell GR, Ross R (1979) The smooth muscle cell in culture. Physiol Rev 59: 1–61.
14. Hall IP, Kotlikoff M (1995) Use of cultured airway myocytes for study of airway remodeling in asthma. Cell. Mol Physiol 291: L281–288.
15. Matsuda F, Sugahara K, Sugita M, Sadohara T, Kiyota T, et al. (2000) Biphasic activation of voltage-dependent calcium channels in airway smooth muscle cells. Am J Physiol Lung Cell Mol Physiol 278: C106–C113.
16. Semenov I, Wang B, Herlihy JT, Brenner R (2011) BK channel beta1 subunits regulate airway contraction secondary to M2 muscarinic acetylcholine receptor mediated depolarization. J Physiol 589: 1803–1817.
17. Liu JQ, Yang D, Felz RJ (2006) A novel bronchial ring bioassay for the evaluation of small airway smooth muscle function in mice. Am J Physiol Lung Cell Mol Physiol 294: L2181–2185.
18. Tomasz M, Boyle JP, Worley JF, III, Kodikoff MJ (1992) Contractile agonists activate voltage-dependent calcium channels in airway smooth muscle cells. Am J Physiol 263: C106–113.
19. Kamishima T, Nelson MT, Patlak JB (1992) Carbachol modulates voltage sensitivity of calcium channels in bronchial smooth muscle of rats. Am J Physiol 263: C69–77.
20. Kajita J, Yamaguchi H (1993) Calcium mobilization by muscarinic cholinergic stimulation in bovine single airway smooth muscle. Am J Physiol 264: L496–503.
21. Matsuda F, Sugahara K, Sugita M, Sadohara T, Kiyota T, et al. (2000) Comparative effect of aramino, aminophylline and diltiazem on rat airway smooth muscle. Acta Anaesthesiolog Scand 44: 763–766.
22. Morey C, Estephan R, Abbott GW, Levi R (2008) Cardioprotective effect of histamine H3-receptor activation: pivotal role of G beta gamma-dependent inhibition of voltage-operated Ca2+ channels. J Pharmacol Exp Ther 326: 871–878.
23. Kinomura T, Kobayashi S, Horiuti K, Somlyo AV, Somlyo AP (1989) Receptor-coupled, permeabilized smooth muscle. Role of the phosphatidylinositol cascade, G-proteins, and modulation of the contractile response to Ca2+. J Biol Chem 264: 3339–3342.
24. Jansen LJ (2002) Ionic mechanisms and Ca2+ regulation in airway smooth muscle contraction: do the data contradict dogma? Am J Physiol Lung Cell Mol Physiol 282: L1161–L1178.
25. Groen R, Zaagman J, Meurs H, Halayko AJ (2006) Muscarinic receptor signaling in the pathophysiology of asthma and COPD. Respir Res 7: 73.
26. Semenov I, Wang B, Herlihy JT, Brenner R (2011) BK channel beta1 subunits regulate airway contraction secondary to M2 muscarinic acetylcholine receptor mediated depolarization. J Physiol 589: 1803–1817.
27. Liu JQ, Yang D, Felz RJ (2006) A novel bronchial ring bioassay for the evaluation of small airway smooth muscle function in mice. Am J Physiol Lung Cell Mol Physiol 294: L2181–2185.
28. Tomasz M, Boyle JP, Worley JF, III, Kodikoff MJ (1992) Contractile agonists activate voltage-dependent calcium channels in airway smooth muscle cells. Am J Physiol 263: C106–113.
29. Kamishima T, Nelson MT, Patlak JB (1992) Carbachol modulates voltage sensitivity of calcium channels in bronchial smooth muscle of rats. Am J Physiol 263: C69–77.
30. Janssen LJ (2002) Ionic mechanisms and Ca2+ regulation in airway smooth muscle contraction: do the data contradict dogma? Am J Physiol Lung Cell Mol Physiol 282: L1161–L1178.
31. Groen R, Zaagman J, Meurs H, Halayko AJ (2006) Muscarinic receptor signaling in the pathophysiology of asthma and COPD. Respir Res 7: 73.
32. Semenov I, Wang B, Herlihy JT, Brenner R (2011) BK channel beta1 subunits regulate airway contraction secondary to M2 muscarinic acetylcholine receptor mediated depolarization. J Physiol 589: 1803–1817.
33. Akhtar S, Mazzeo AJ, Cheng EY, Bosnjak Z, Kampine JP (1999) Differential contributions: Edited paper: LML. Conceived and designed the experiments: RZG. Performed the experiments: CZ RZG. Analyzed the data: CZ LML KEF RZG. Contributed reagents/materials/analysis tools: KFU MEM. Wrote the paper: RZG.

**Acknowledgments**

We thank Walter H. Hsu at Iowa State University for helpful discussions, the staff in the Department of Pathology at the University of Massachusetts Memorial Medical Center for help with collecting human lung tissue, and Keita Shibata for technical assistance in force measurement. CZ expresses his gratitude to Min-sheng Zhu at Nanjing University for his support to study at the University of Massachusetts Medical School.

**Author Contributions**

The author(s) have made the following declarations about their contributions: Edited paper: LML. Conceived and designed the experiments: RZG. Performed the experiments: CZ RZG. Analyzed the data: CZ LML KEF RZG. Contributed reagents/materials/analysis tools: KFU MEM. Wrote the paper: RZG.

**Movie S1** This clip shows 1 mM chloro reversed the 100 \(\mu\)M Mch-induced increase in [Ca\(^{2+}\)], and cell shortening; the first 120 images of this clip were analyzed and plotted in Figure 2A. The images are displayed as fluorescence intensity (rather than \(\Delta F/F_0\) because the cell changes its shape dramatically in response to stimuli (and changing thickness makes \(\Delta F/F_0\) measures misleading).

**MOV**
idene Ca2+ channel activator, in cardiac and vascular preparations. Mol Pharmacol 40: 734–741.

40. Murlas C, Ehring G, Sznitko J, Sperelakis N (1986) Kr+-induced alterations in airway muscle responsiveness to electric field stimulation. J Appl Physiol 61: 60–67.

41. Mitchell RW, Murphy TM, Kelly E, Leff AR (1991) Extracellular Ca2+ mobilization in potential-dependent contraction of tracheal smooth of maturing snake. J Appl Physiol 71: 1489–1495.

42. Farley JM, Miles PR (1957) Role of depolarization in acetylcholine-induced contractions of dog tracheal muscle. J Pharm Exp Ther 201: 199–205.

43. Coburn RF (1979) Electromechanical coupling in canine tracheal muscle: acetylcholine contractions. Am J Physiol 236: C177–184.

44. Daenas C, Hatziefthimiou AA, Gourgoulian KI, Molyvdas PA (2006) Azithromycin has a direct relaxant effect on precontracted airway smooth muscle. Eur J Pharmacol 553: 280–287.

45. Liu C, Zuo J, Pertens E, Helli PB, Janssen LJ (2005) Regulation of Rho/ROCK signaling in airway smooth muscle by membrane potential and [Ca2+]i. Am J Physiol Lung Cell Mol Physiol 289: L574–582.

46. Mbikou P, Fujaut M, Bruinen M, Roux E (2011) Contribution of Rho kinase to paradoxical increase in intracellular Ca2+ in the early phase of the calcium-contraction coupling in airway smooth muscle. Exp Physiol 96: 240–254.

47. Perez JF, Sanderson MJ (2005) The frequency of calcium oscillations induced by 5-HT, ACH, and KCl determine the contraction of smooth muscle cells of intrapulmonary bronchi. J Gen Physiol 125: 535–553.

48. Janssen LJ, Tazewo T, Zuo J, Pertens E, Kesquaree S (2004) KCl evokes contraction of airway smooth muscle via activation of RhoA and Rho-kinase. Am J Physiol Lung Cell Mol Physiol 287: 1452–1458.

49. Hirota S, Janssen LJ (2007) Store-refilling involves both L-type calcium channels and a reverse-mode sodium-calcium exchanger in airway smooth muscle. Eur Respir J 30: 269–278.

50. Wu SV, Chen MC, Rozengurt E (2005) Genomic organization, expression, and function of bitter taste receptors (T2R) in mouse and rat. Physiol Genomics 22: 179–193.

51. Shah AS, Ben-Shahar Y, Moninger TO, Kline JW, McNich MJ (2009) Motile cilium of human airway epithelium are chemosensitive. Science 325: 1111–1114.

52. Tizzano M, Gulbransen BD, Vandenbeuch A, Clapp TR, Herman JP, et al. (2011) Cholinergic chemosensory cells in the trachea regulate breathing. Proc Natl Acad Sci U S A 108: 30143–30151.

53. Patel KR, Tullett WM (1985) Comparison of two calcium antagonists, verapamil and diltiazem on airway reactivity to methacholine and exercise in subjects with mild intermittent asthma. Am Rev Respir Dis 136: 1179–1182.

54. Ikeda SR (1996) Voltage-dependent modulation of N-type calcium channels by G-protein beta gamma subunits. Nature 380: 255–258.

55. Xu J, Cao J, Iguchi N, Riethmacher D, Huang L (2012) Functional and protein-gated inhibitory module of N-type (ca(v)2.2) ca2+ channels. Nature 480: 891–904.

56. Kohlkoff MI (1980) Calcium currents in isolated canine airway smooth muscle cells. Proc Natl Acad Sci U S A 108: 9478–9483.

57. Pietras CO, James A, Konradsen JR, Nordlund B, Soderhall C, et al. (2012) Intracellular Ca2+ silences L-type Ca2+ channels in mesenteric veins: inhibition of venous smooth muscle resistance to calcium channel blockers. Circ Res 106: 739–747.

58. Meyerhof W, Batram C, Kuhn C, Brockhoff A, Chudoba E, et al. (2010) The mechanism of venous smooth muscle resistance to calcium channel blockers. Circ Res 106: 739–747.

59. Cheng H, Yibchok-Anun S, Park SC, Hsu WH (2002) Somatostatin-induced calcium channel activator, in cardiac and vascular preparations. Mol Pharmacol 40: 734–741.

60. Thakali KM, Kharade SV, Sivasubramanian S, Altered B, et al. (2010) In vivo superresolution three-dimensional images of fluorescence in cells with minimal light exposure. Science 268: 1483–1487.

61. Brockhoff A, Behrens M, Massarotti A, Appendino G, Meyerhof W (2007) Regulation of vascular L-type Ca2+ channels by phosphatidylinositol 3,4,5-triphosphate. Circ Res 95: 300–307.

62. Nino G, Hu A, Grunstein JS, McDonough J, Kreiger PA, et al. (2012) G Protein beta-gamma subunit signaling mediates airway hyperresponsiveness and inflammation in allergic asthma. PLoS ONE 7: e32078. doi:10.1371/journal.pone.0032078

63. Huang L, Shanker YG, Dubaukaitse J, Zheng JZ, Yan W, et al. (1999) Ggum13 colocalizes with gustducin in taste receptor cells and mediates IP3 responses to bitter androgenous tastes. J Neurosci 2: 1053–1062.

64. Kreisler G, Canning BJ, Hartmann P, Veres TZ, Papadakis T, et al. (2011) Syntillas C, Ehring G, Sznitko J, Sperelakis N (1986) Kr+-induced alterations in airway muscle responsiveness to electric field stimulation. J Appl Physiol 61: 60–67.

65. Liu C, Zuo J, Pertens E, Helli PB, Janssen LJ (2005) Regulation of Rho/ROCK signaling in airway smooth muscle by membrane potential and [Ca2+]i. Am J Physiol Lung Cell Mol Physiol 289: L574–582.

66. Liu C, Zuo J, Pertens E, Helli PB, Janssen LJ (2005) Regulation of Rho/ROCK signaling in airway smooth muscle by membrane potential and [Ca2+]i. Am J Physiol Lung Cell Mol Physiol 289: L574–582.

67. Liu C, Zuo J, Pertens E, Helli PB, Janssen LJ (2005) Regulation of Rho/ROCK signaling in airway smooth muscle by membrane potential and [Ca2+]i. Am J Physiol Lung Cell Mol Physiol 289: L574–582.

68. Thakali KM, Kharade SV, Sivasubramanian S, Altered B, et al. (2010) In vivo superresolution three-dimensional images of fluorescence in cells with minimal light exposure. Science 268: 1483–1487.

69. Weiss N, Lenigrand C, Pourraux S, Bicharouaa H, Allard B, et al. (2010) In vivo superresolution three-dimensional images of fluorescence in cells with minimal light exposure. Science 268: 1483–1487.