Intragastric nitric oxide production in humans: measurements in expelled air

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Abstract
High values (800–6000 parts per billion) of nitric oxide (NO) in expelled air from the stomach were shown in humans by chemiluminescence technique. These NO values were more than 100 times higher than those found in orally exhaled air. Intragastric NO production is probably non-enzymatic, requiring an acidic environment, as NO in expelled air was reduced by 95% after pretreatment with the proton pump inhibitor omeprazole. Furthermore, large amounts of NO were formed in vitro from lettuce and saliva when placed in hydrogen chloride (pH <2). In conclusion, large amounts of NO are formed intragastrically in humans and this source of NO may be of importance for the integrity of the gastric mucosa in health and disease. Measurements of NO in expelled air might be of value as a non-invasive method for estimation of gastric acidity.

Human saliva contains both nitrate and nitrite.1 Nitrate is absorbed in the gastrointestinal tract and extracted from plasma into saliva by the salivary glands.1 Most of the nitrite in saliva results from reduction of nitrate by a variety of micro-organisms in the oral cavity.1 Regularly swallowed saliva will continuously enter an acidic environment in the stomach, and under these conditions nitrite may be reduced to nitric oxide (NO).2 We wanted to investigate if NO is formed intragastrically in humans and to compare that with NO formation from saliva and nitrate containing food in an acidic environment in vitro. Furthermore, as increased values of NO have recently been shown in exhaled air of asthmatic subjects,1 we also wanted to examine if NO formation in the stomach would influence the values of NO in exhaled air. Parts of these results have been presented in a preliminary form.4

Methods
SUBJECTS
Four healthy non-smoking subjects (29–40 years old) and four non-smoking atopic subjects (30–40 years old), with confirmed allergy towards at least rat allergen, and symptoms of mild asthma and rhinitis took part in the study. One of the asthmatic subjects inhaled a glucocorticoid (budesonide) regularly and the other three inhaled a β2 agonist or sodium cromoglycate when necessary. All subjects were tested when they were subjectively free from respiratory tract infections and none reported any symptoms of dyspepsia. The procedures used in this study have been reviewed and approved by the local ethical committee.

NITRIC OXIDE VALUES IN EXPELLED AIR FROM THE STOMACH
Voluntary regurgitation of air was performed three to five minutes after intake of 30 cl of carbonated water with a pH of 5.5 (Ramlösa; Pripps AB, Sweden). Expelled air was led into a Teflon tubing system from which air was continuously sampled (0.8 l/min) into an NO chemiluminescence analyser (CLD 700; Eco Physics, Dürnten, Switzerland), and peak concentrations of NO were registered on a chart recorder. The detection limit for NO was 1 part per billion (ppb) and calibrations at known concentrations of NO in N2, using an electromagnetic flow controller (Environics Inc, Middletown CT, US) were performed. The chemiluminescence assay is highly specific for NO and does not interfere with other nitrogen oxides.3 Measurements of NO in expelled air from the stomach were made after 10 hours of fasting in combination with one of the following pretreatment procedures: (1) no pretreatment (control); (2) intake of 50 g of iceberg lettuce (nitrate load). The mean nitrate content in ordinary iceberg lettuce is 13 g/kg; (3) pretreatment by oral intake of a total of 240 mg (three times 80 mg) of the proton pump inhibitor omeprazole (Astra-Hässl AB, Göteborg, Sweden) distributed over a 24 hour period before the experiments; (4) intake of 50 g of lettuce after omeprazole pretreatment.

NITRIC OXIDE FORMATION IN VITRO
A glass bottle (1 litre) standing on a heat plate with a magnetic stirrer was used. The investigated test substances were placed in the glass bottle, which was then closed during an incubation time of five minutes. The substances used were: 30 cl of carbonated water in 50 ml of HCl (100 mM, pH 1.0); 50 g of lettuce (cut in small pieces) in 50 ml of HCl (100 mM, pH 1.0); 6–8 g of saliva in 50 ml of HCl (100 mM, pH 1.0), with and without 30 cl of carbonated water; 2 g of saliva collected after 10 hours of fasting or two hours after intake of 250 g of lettuce, in 50 ml of HCl (100 mM, pH 1.0); 50 g of chewed lettuce, containing 6–8 g of saliva in 50 ml of different HCl solutions with pH ranging from 0.5 to 4.0 (316–0.1 mM) (saliva content was calculated...
NITRIC OXIDE VALUES IN EXHALED AIR

The subjects wore a nose clip and breathed NO free air (NO < 2 ppb; AGA AB, Lidingö, Sweden) with normal tidal volumes through a mouth piece connected to a non-rebreathing valve. Exhaled air was led into a Teflon tubing system from which air was continuously sampled (0.8 l/min), and steady state concentrations of NO during continuous breathing were recorded. NO measurements were made after 10 hours of fasting with and without omeprazole pretreatment.

STATISTICS

All data are presented as mean (SEM). Wilcoxon's signed rank test and Mann-Whitney U test were used for paired and unpaired comparisons, respectively.

Results

NITRIC OXIDE VALUES IN EXPELLED AIR FROM THE STOMACH

Control NO concentrations in expelled air after 10 hours of fasting were 602 (107) ppb and these values increased fourfold, five minutes after intake of lettuce (Fig 1). Pretreatment with omeprazole considerably reduced the NO concentrations in expelled air both without (96%) and with (95%) intake of lettuce.

NITRIC OXIDE FORMATION IN VITRO

NO formation was strongly pH dependent between pH 1 to 2, when incubating chewed lettuce (containing 6-8 g of saliva) in HCl (Fig 2). The values at pH 1 were 25 times higher than those formed from saliva alone in HCl and 500 times higher than from lettuce alone in HCl. The addition of 30 cl of carbonated water reduced NO formation by about 50% (Fig 3). Carbonated water alone in HCl did not produce any significant amounts of NO. In a separate experiment, saliva collected two hours after intake of 250 g of lettuce and placed in HCl (pH 1-0), yielded five times higher NO values than saliva collected after 10 hours of fasting (1640 (420) ppb and 334 (110) ppb respectively, p<0.05). NO formation was also pH dependent when incubating a nitrite solution in HCl at different pH (not shown). At pH 1-0, NO concentrations from the nitrite solution and saliva were similar (Fig 3). Nitrate solutions did not yield any NO when placed in HCl at pH 1-0.

Discussion

In this study we have identified the stomach as...
an important source of NO at values greatly exceeding those in orally exhaled air. Gastric NO formation seems to be dependent on an acidic environment, as inhibition of gastric acid secretion by the selective proton pump inhibitor omeprazole almost abolished NO in expelled air. Furthermore, lettuce and saliva in HCl in vitro, yielded high NO concentrations at pH 1, with a gradual decrease at higher pH. NO formation from chewed lettuce in HCl was pH dependent with very low concentrations of NO formed at a pH > 3, a pH probably obtained in the stomach in vivo after omeprazole pretreatment. The in vitro model with 50 ml of HCl, pH 1·0 was chosen as this resembles the physiological situation regarding the amount and acidity of gastric juice during fasting conditions.

The substrate for intragastric NO formation is probably nitrite, as it is well known that nitrite is reduced in an acidic environment, thus forming NO.2 Gastric juice normally contains small but significant amounts of nitrite,11 which may be derived from swallowed saliva containing both nitrate and nitrite,1 or possibly from active gastric secretion of nitrite.11 Most of the nitrite in saliva is formed through the reduction of salivary nitrite by bacteria in the oral cavity.1 The daily saliva production in humans often exceeds one litre, thus permitting continuous formation of NO in the stomach over the day. These suggestions are supported by the fact that saliva alone in HCl in vitro yielded high amounts of NO, similar to those produced from a nitrite solution in HCl. Furthermore, saliva collected after a nitrate load produced five times more NO compared with saliva collected after 10 hours of fasting when placed in HCl. These results are in accordance with earlier studies showing that the nitrite concentration in saliva reaches a maximum two hours after a nitrate load.1

Intake of nitrate rich food (lettuce) increased the NO values in expelled air fourfold compared with fasting conditions. This could be explained by the high amounts of nitrate in lettuce2 being reduced to nitrite by bacterial enzymes in the saliva, which is also supported by the finding that chewed lettuce in HCl in vitro formed much more NO than saliva alone in HCl. An alternative explanation may be that agents with reducing capacity in the lettuce could enhance nitrite formation from nitrate in the saliva.

As the stomach is anatomically related to the airways through the oesophagus, we also wanted to investigate its possible contribution to NO concentrations in exhaled air. The values of NO in expelled air after fasting were 100–1000 times higher than those in orally exhaled air. Hence, only minute amounts of expelled air may influence the exhaled values bearing in mind that asthmatic subjects may have a high incidence of gastro-oesophageal reflux.12 The high NO concentrations in the stomach, however, did not seem to contribute continuously to NO concentration in exhaled air during normal breathing in healthy subjects or asthmatic subjects, as these values were not affected by omeprazole pretreatment.

NO measurements in expelled air after a standard ingestion of nitrite could possibly be of clinical use, for example, in the diagnosis of achlorhydria and in monitoring of peptic ulcer disease, as it may reflect gastric acidity.

The finding that the stomach is an important source of NO is interesting, as NO may be important in bacteriostasis13 and in the regulation of mucosal blood flow, mucus formation,14 and possibly gastric motility including lower oesophageal sphincter tonus.15 Instillation of the NO donor isosorbide dinitrate into the rat gastric lumen in vivo produced an increase in mucus formation that was prevented by coadministration of oxyhaemoglobin, suggesting that the action was mediated by local release of NO.16 Furthermore, oral administration of nitrates in the rat prevents the formation of HCl induced gastric lesions, and topical application of nitrates to the gastric mucosa increases mucosal blood volume.17 Both these studies show that luminal NO may affect the gastric mucosa. The possible influence of the comparatively high amount of NO in the gastric lumen, even during fasting conditions, on the development of diseases such as peptic ulcer18 and stomach cancer19 needs further study.

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