Neuropeptides of the autonomic nervous system in Sjögren’s syndrome

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Abstract

Objective—To assess the activity level of the autonomic nervous system in Sjögren’s syndrome (SS) and to correlate this with stress.

Methods—Patients with SS (n=12) and healthy controls (n=10) were analysed for the content of vasoactive intestinal peptide (VIP) and neuropeptide Y (NPY) in their stimulated saliva by radioimmunoassays and for stress by the use of a modified Jenkins Activity Survey (JAS).

Results—The data are expressed as median (interquartile range). Salivary VIP output (pg/min) and NPY output (pg/min) were high in SS compared with healthy controls (30.0 (15.6, 36.6) versus 12.3 (9.2, 24.0), p<0.045, 4.8 (0.6, 24.1) versus 0.7 (0.0, 2.4), p=0.038, respectively). Patients experienced only a little, but not significantly, more stress than the healthy controls (stress index −2.8 (−7.7, 6.9) versus −5.2 (−12.9, 2.7), p>0.05). Stress in general was associated with high salivary VIP concentrations (r=0.41, p=0.05).

Conclusions—These findings show that adequately processed saliva (containing aprotinin and EDTA as neuropeptidase inhibitors) contains measurable amounts of marker peptides of the autonomic nervous system. Secondly, VIP concentration but not output may be affected by stress, which may act by decreasing salivary flow. In patients with SS, VIP and NPY outputs are increased. This may indicate increased leakage into saliva or efforts to compensate for the diminished salivary flow, or both.

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Sjögren’s syndrome (SS) is a systemic autoimmune disease characterised by exocrinopathy of unknown aetiology and pathogenesis leading to dry mouth (xerostomia) and dry eyes (keratoconjunctivitis sicca). A particularly prominent and diagnostically useful feature is focal adenitis which develops in both salivary and lacrimal glands. For practical reasons focal adenitis is usually diagnosed from labial salivary gland biopsy specimens. This focal sialadenitis is associated with decreased resting salivary flow (according to the European criteria ≤0.1 ml/min). There is a remarkable discrepancy between the focal and often relatively mild glandular involvement in form of adenitis (with pathological focus score values ≥1 often affecting less than 10% of the total area, except in far advanced end stage cases) and the considerably decreased salivary flow (≤0.1 ml/ml of whole resting salivary flow—that is, about only 10% of the normal salivary production). One explanation, which has recently been put forward, is that this autoimmune disorder as such or as a result of focal adenitis leads to vasoneural dysregulation and injury of the peripheral nerve fibres and thus to diminished salivary flow and atrophy of the acinar cells. It has been widely accepted that normal salivary flow is under neural regulation: parasympathetic stimulation increases salivary flow, whereas stressful stimuli decrease it via central sympathetic activation. Accordingly, a parasympathomimetic drug, pilocarpine, has been found useful in the treatment of dry mouth and electrostimulation, which augments normal physiological salivary reflexes, stimulates production of saliva in SS.

The aim of the study was therefore to measure the salivary content of vasoactive intestinal peptide (VIP) and neuropeptide Y (NPY), which in labial salivary glands are found mainly in the parasympathetic and sympathetic arm, respectively, of the autonomic nervous system. Activity of the autonomic nervous system is influenced by the somatic well being, in particular by the degree of stress. In fact, dry mouth associated with depression and anxiety or the treatment with psychotropic drugs (with anticholinergic side effects), or both, is one of the most important differential diagnostic alternatives in Sjögren’s syndrome clinics. We therefore also aimed to measure stress by using a modified Jenkins Activity Survey (JAS) to correlate it with the degree of sympathetic and parasympathetic activity.

Methods

PATIENTS AND CONTROLS

Twelve female patients with chronic established SS were studied; all fulfilled the European Classification Criteria for SS. Primary SS was diagnosed if any four of the six classification criteria are fulfilled. Secondary SS is diagnosed if there is a positive response to items 1 or 2 (ocular symptoms or oral symptoms), plus a positive response to at least two items from among 3, 4, and 5 (ocular signs, histopathological features, salivary gland involvement). In both cases, the adequate exclusions (sarcoidosis, lymphoma, acquired immunodeficiency syndrome and graft versus host disease) were done. Ten healthy female volunteers served as controls.

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The average age of the SS patients was 47.5 (range 21–59) years. The duration of the SS was on average 8.6 (range 1–17) years. All the patients were non-smokers. Five of the patients were married, three unmarried, and four divorced. Three of them worked in a white collar profession, four in a blue collar profession, one was a college student, and four were retired because of the disease.

The average age of the 10 healthy controls was 47.7 (range 39–56) years. Six of them were married, one was engaged to be married, two were divorced, and one was a widow. Nine of the controls were blue collar workers in a health care profession and one was a cleaning woman. Eight of them were non-smokers, one smoked 5–20 cigarettes a day, and another smoked more than 20 cigarettes.

RADIOIMMUNOASSAY
Both the patients and the controls gave a stimulated salivary sample as described elsewhere. The stimulated whole saliva was collected for 15 minutes. The stimulus used was chewing a piece of hard paraffin. The saliva samples were collected to chilled tubes containing 50 000 IU/ml aprotinin and 10 mM EDTA to minimise artefactual neuropeptide degradation in vitro and stored at −40°C until analysed.

Neuropeptide concentration in saliva was measured using a standard competitive 125I-radioimmunoassay kit (Peninsula, Belmont, CA, USA). Briefly, (1) 100 µl of sample or standard was incubated in polypropylene tubes for 24 hours at +4°C with 100 µl rabbit antihuman VIP or NPY IgG, (2) 100 µl of 125I-labelled peptide (10 000–15 000 cpm/100 µl) was added and incubated for 24 hours at +4°C; (3) 100 µl goat antirabbit IgG and 100 µl normal rabbit serum were added and incubated for 90 minutes at room temperature before (4) 500 µl RIA buffer was added followed by centrifugation at 1700 g for 20 minutes at +4°C; supernatant was disregarded and the radioactivity of the pellet was measured with a programmed gamma scintillation counter. Rabbit antihuman VIP antibodies show no cross reactivity with human PHM-27 (peptide histidine methionine), substance P, VIP 1-12, endothelin-1, glucagon, somatostatin, PACAP-38 (pituitary adenylate cyclase activating peptide), PACAP-38 (16-38), porcine secretin or rat galanin and less than 0.02% cross reactivity with human PACAP-27. Similarly, rabbit antihuman NPY antibodies show no cross reactivity with human pancreatic polypeptide, VIP, amylin, prepro NPY 68-97, insulin or somatostatin. All samples were analysed in duplicate with individual controls for apparent binding in the absence of antibody (that is, total counts, nonspecific binding, total binding).

JENKINS ACTIVITY SURVEY
The JAS scale consists of 84 items that are divided in four subscales. For this study we recorded the values of the actual A-B factor with 21 items, which best reflect the stress behaviour of a person.

The study was carried out according to the principles of the Declaration of Helsinki, and with the approval of the ethical committees of the institutions involved. The BMDP software package version 7.0, was used for the statistical analysis. The data are expressed as median (interquartile range). The comparison between the patients and the healthy controls was performed with the use of the Mann-Whitney test. The difference in variability between samples was tested by Siegel-Tukey test when there was no difference in median and by Moses rank-like test when there was a difference. The correlation between stress and salivary neuropeptide content was studied with the use of the Spearman’s rank correlation coefficient. The reliability of the stress factor was measured with the use of the Cronbach alpha test.

Results
SALIVARY FLOW RATE
There was no significant difference in the median between the stimulated flow rate (ml/min) of the SS patients and the healthy controls (0.4 (0.2–0.7) v 0.4 (0.3–0.4), p=0.304), but there was a significant difference in the variability between the flow rate values of the groups (p = 0.0018) (fig 1).

SALIVARY VIP AND NPY
Salivary VIP output (pg/min) was high in SS compared with healthy controls (33.6 (22.6–64.8), p=0.025). There was no significant difference when expressed per mg protein or content (22.4 (18.1–30.6) v 15.2 (7.7–38.2), p>0.05). The concentration and content of salivary VIP in the two smokers were numerically below the average of the other healthy controls.

Salivary NPY output (pg/min) was high in SS compared with healthy controls (4.8 (0.6–24.1) v 0.7 (0.0–2.4), p=0.038), also there was significant difference in variability (p=0.016) (fig 3). Salivary NPY concentration (3.4 (1.2–7.6) pg/ml v 2.1 (0.0–7.1) pg/ml)
and content (pg/mg protein: 2.3 (0.6–27.3) vs. 2.7 (1.6–4.4)) did not significantly differ in SS and in healthy controls and there were no differences in variability. NPY contents in the two smokers were similar to NPY values in other healthy controls.

**Discussion**

SS is characterised by sialopenia/xerostomia. It is the resting salivary flow, however, which is low. Stimulated salivary flow varies a lot between patients and may be normal. It has therefore not been included in the European Classification criteria. As a matter of fact, many patients use sialogogic stimuli, mechanical (such as a xylitol chewing gum) or chemical (such as pilocarpin) to alleviate their symptoms. In advanced or difficult cases, or both, stimulated salivary flow/reserves are low, but in many series including newly diagnosed patients stimulated salivary flow is normal. This was also the case in our series.

VIP is a 28 amino acid peptide, which belongs to the family of brain-gut peptides. VIP in salivary glands it often colocalises with choline acetyltransferase and seems therefore to be useful as a peptide marker for postganglionic parasympathetic fibres in that tissue. NPY is a 36 amino acid peptide of the NPY/CPON family. It usually colocalises with cytoplasmic tyrosine hydroxylase and intravesicular dopamine-â-hydroxylase, which are classic markers for postganglionic sympathetic nerves. Both VIP and NPY containing peptidergic nerves have been recently described in salivary glands in normal healthy controls and in patients with SS.

In this study the eventual release into and presence in saliva of VIP and NPY was assessed by RIA from saliva collected to a cocktail of enzyme inhibitors selected to prevent artefactual in vitro degradation by various neuropeptidases. Both neuropeptides were found in the whole series, stress score (as expressed by the JAS factor A-B type behavior) correlated significantly with both VIP concentration (pg/ml, \( r=0.41, p<0.05 \)) and content (pg/mg protein, \( r=0.53, p=0.02 \)). Thus, the more stress, the higher the VIP values. The correlation between NPY concentration or content and stress did not quite reach significance (\( r=0.25, p>0.05; r=0.33, p>0.05 \)).

When all study subjects were subdivided into patients and controls, it was found that in the patient group there were no significant correlations between the stress score and VIP values in the saliva, although there was a significant correlation between the stress score and the NPY concentration (\( r=0.66, p=0.02 \)). There were positive, but not statistically significant, correlations in the healthy controls between the stress score and the VIP content and the VIP output (\( r=0.52, p>0.05; r=0.54, p>0.05 \) respectively).

**JAS**

The reliability of the type A-B factor was tested with the use of the Cronbach alpha coefficient (alpha = 0.51) and on basis of this calculation, five of the original 21 items were omitted. After omission of these factors, the alpha coefficient improved to 0.63. However, also when the corrected mean value for the A-B type factor was used, the patient group (−2.8 (−7.7, 6.9)) did not differ significantly from that of the control group (−5.2 (−12.9, 2.7); \( p>0.05 \)). The duration of the patients’ disease history, the severity of the general disease symptoms (joint pain and fatigue), the severity of local symptoms (symptoms of the eyes, dryness of the mouth), and the patients’ age were not significantly related to the value of the A-B factor.

**CORRELATION BETWEEN SALIVARY NEUROPEPTIDES AND STRESS**

In the whole material, there were no significant correlations between stress (as expressed by the JAS factor A-B type behaviour) and the output of the neuropeptides. In the whole series, stress score (as expressed by the JAS factor A-B type behavior) correlated significantly with both VIP concentration (pg/ml, \( r=0.41, p=0.05 \)) and content (pg/mg protein, \( r=0.53, p=0.02 \)). Thus, the more stress, the higher the VIP values. The correlation between NPY concentration or content and stress did not quite reach significance (\( r=0.25, p>0.05; r=0.33, p>0.05 \)).

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adequately processed samples suggesting local release and secretion to saliva. After this first finding on the presence of VIP and NPY in saliva, eventual effects of disease and stress on their concentration were evaluated. These are often connected, because psychosomatic features associated with chronic diseases contribute to psychic and somatic well-being. The JAS instrument describes the person’s stress behaviour. The type A behavioural pattern comprises a cluster of actions and emotions exhibited by people who engage themselves in a chronic struggle and have a hyperresponsiveness to almost any challenge. These people are competitive with a driving need for control and an accentuated sense of time urgency. Changes in life style or circumstances as well as disease are known to modify the person’s type A behavioural pattern. While stress may affect health, there is also evidence that a state of ill health, as was the case with our SS patients, can act as a significant source of stress in itself or can sensitize the person to other sources of stress by reducing their ability to cope. Furthermore, at any point of time, the relation between stress and disease can operate in both directions, and may develop into a vicious cycle.

SS patients experienced on average slightly but not significantly more stress than the healthy controls as measured with the JAS instrument. According to the present findings, patients with SS seem to cope surprisingly well with their disease, at least as assessed by the extent of the stress they experience.

The physiological factors regulating VIP and NPY secretion are not known. In this study we found an interesting correlation between stress and salivary VIP concentration and content, but not VIP output. Stress might increase VIP release or, more probably, decrease the watery salivary flow. Stress may interfere with the normal salivary gland function finely controlled by coordinated and balanced excitatory and inhibitory inputs from tonically active postganglionic neurons. VIP has many different paracrine effects. It induces vasodilatation, modulates mast cell mediator release and lymphocyte migration and recirculation. In addition, VIP has important trophic effects on salivary gland acinar cells in the long term. According to recent immunoenzyme microscopic studies published elsewhere, VIPergic fibres come into close contact with the acinar cells and some fibres are even located hypoelemally—that is, inside the acinar basement membrane. NPY causes vasoconstriction, modulates neurotransmitter release from sensory and postganglionic sympathetic neurons, and up regulates adhesiveness of endothelial cells for leucocytes. According to this study, these neuropeptides are not only found in salivary glands, but are also released there and to some extent found in saliva; they might therefore contribute to the local pathomechanisms in SS.

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