Nasal Air-Conditioning During Breathing Therapy

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Running Title:
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ABSTRACT

It has been reported that continuous positive airway pressure therapy introduces negative nasal side-effects including sneezing, itching, nasal dryness, nasal congestion and/or a runny nose. As these symptoms are suggestive of nasal dysfunction, heated humidification is often used to fully saturate and heat the inhaled air to core body temperature. It is expected that this relieves the nasal mucosa from having to supply, or recover, heat and moisture from inspired and expired air. This review summarizes the current in vitro and in vivo knowledge relevant to nasal air-conditioning, and identifies further investigations necessary to improve our understanding the changes that occur during nasal continuous positive airway pressure therapy. Investigations into nasal airway fluid transportation, airflow regulation and heat and fluid supply may lead to a therapy temperature/pressure/humidification algorithm that optimizes these parameters for a prescribed therapy pressure. Optimization could lead to a reduction in titration pressure and improved treatment compliance.

Keywords

nasal air-conditioning, nasal mask, breathing therapy.
INTRODUCTION

The nasal mask is a common interface through which air or other gas mixtures are delivered to a spontaneously breathing patient. Breathing dry gases through a nasal mask presents significant air-conditioning challenges to the nose which may lead to desiccation of the nasal mucosa. Heated pass-over type humidifiers are commonly used to condition the gas to near full water saturation at core body temperature [1-3]. Other elevated pressure breathing therapy applications, such as nasal continuous positive airway pressure (n-CPAP) treatment for obstructive sleep apnea (OSA), utilize similar supplementary heating and humidification. Pressure related nasal complications have been reported by over 40% of n-CPAP users [4, 5]. These include nasal crusting, dryness, congestion, sneezing, rhinorrhea and/or itching. The nose appears unable to adequately condition inhaled ambient air under slightly elevated pressures [2-4, 6-12]. Heated humidification is commonly utilized to treat these symptoms and improve patient comfort; however, its ability to improve treatment compliance is questionable [5, 7, 13].

Currently no clear explanation exists as to why breathing ambient air at slightly elevated pressures induces these nasal symptoms. It has been suggested that these symptoms could have existed prior to initiation of therapy, however they commonly disappear with cessation of treatment [9]. Another suggestion attributes these symptoms to mouth leak [1, 14], where large unidirectional airflow occurs through the nose. This type of flow prevents expired air heat and moisture recovery from occurring within the airway fluid lining. These symptoms are also attributed to changes in nasal blood flow which occur due to the elevated pressures [1, 15]. Further, age can contribute to negative nasal symptoms during breathing therapy. Functional changes occurring within the nose such as mucosal atrophy [8, 16], rather than nasal mask conditions, may also be responsible for these symptoms.

Within the airways, the ciliated respiratory epithelium is protected by a superficial gel layer of mucus, which is transported along the surface by the synchronized beating of the cilia within the periciliary liquid (PCL) or sol layer. These two fluid layers make up the airway surface liquid (ASL) lining the entire airway. Currently, heating and humidification requirements within most breathing therapy systems are pre-set in terms of both temperature and moisture content of the air delivered to the patient [5, 17]. This conservative approach provides supplementary heated humidification that effectively relieves the nose of any air-conditioning role. Excessive supplementary water will cause dilution of the ASL, whilst
insufficient water can lead to mucus thickening and drying [18]. Both of these conditions adversely affect
mucociliary transport velocity (MTV), heat/moisture exchange and consequently ASL changes, which
could lead to respiratory complications and thermal damage to airway tissue [19, 20].

This paper presents a comprehensive literature review to establish relationships between nasal applied
breathing therapy and symptoms suggestive of airway mucosal dysfunction. This review starts with the
nasal heat and moisture exchange, which is essential for any air-conditioning process, followed by nasal
regulation systems. The latter have been reviewed in the context of their response to the physical
properties of air pressure, flow, humidity and temperature. From a biomedical engineering perspectives,
modeling is an essential part in understanding and predicting any physiological phenomena, modeling is
reviewed followed by air-conditioning pathophysiology. To complete the review, given some of these
relationships have not yet been established yet, ideas for further research are proposed.

NASAL HEAT AND MOISTURE EXCHANGE

Within the nose, there are interactions between independently regulated systems (Fig. 1) in order to
control nasal air-conditioning. Nasal air-conditioning requires heating and water mass transport systems
operating in conjunction with an efficient mucus propulsion system. Regulation of these two fluxes is
achieved through the combined actions of the nasal autonomic nervous system and epithelial cell fluid
transport. Over a 24 hour rest period, healthy humans normally breathe around 10,000 liters of air which
is typically supplemented with around 400 ml of water and 1470 J of heat energy when breathing ambient
air at 25°C temperature and 50% relative humidity [21, 22]. During quiet breathing, in vivo measurements
across the nasal cavity show increased air temperature and humidity levels in an anterior to posterior
direction during both inhalation and exhalation [22].

The nasal fluid lining plays an important role in terms of water and heat transfer during the breathing
cycle [23]. The temperate inspired air, aided by turbulence, is heated and humidified as it passes through
the nasal cavity, causing the mucosa to cool when water transfers from the airway fluid to the air.
However, during exhalation, some of the moisture from the fully saturated air at core body temperature
condenses on the nasal mucosa which has been cooled during inhalation. Exhaled moisture is captured for
use in humidification during the next breathing cycle. The cyclic exchange of latent and sensible heats as
well as the partial water recovery implies that the nasal fluid lining acts as a buffer zone between the nasal
air and the underlying epithelial layer supplying water and heat. This buffering effect possibly reduces the
transient fluid requirement from secretion glands and stabilizes cyclic thermal fluctuations during normal
breathing. It may also offer protection from hot air damage due to the fact that it has a higher sensible
heat capacity than air [18]. The nasal airway fluid lining also provides a platform for mucus transport [24-
26], particulate entrapment and absorption of gaseous water soluble air contaminants [27, 28].

Mucociliary transport of airway fluid is achieved by the synchronized beating of cilia protruding from
pseudo-stratified columnar epithelial cells located in the airway mucosa. Cilia, normally beating around
10-20 Hz in the thinner 5-7 µm thick PCL layer, engage the thicker gel layer [29] with a whip-like motion
[30-33]. The primary function of the upper mucus gel layer is particulate entrapment. Over time this layer
progressively forms into globules, which coalesce into small flakes and subsequently larger plaques.
Continuous transportation of the mucus layer towards the nasopharynx, where it is cleared by swallowing
or expectoration, is essential not only for removal of entrapped contaminants [28, 34], but also for nasal
air-conditioning [18]. Insufficient MTV may lead to an increase in airway fluid depth or a build-up of
coalesced mucus hindering both heat and water exchange. On the other hand, excess airway surface liquid
transported from the lower lungs is absorbed by the epithelial cells [34].

Airflow regimes also play a significant role in heat and water mass transport. The nasal valve region,
anterior to the inferior turbinate, provides a major dynamic portion of nasal airflow resistance. This region
is thought to create turbulent airflow, which enhances heat and water mass transportation [35, 36].
Widening of the air passage posterior to the turbinates with subsequent slowing of airflow enables
deposition of particulate matter. Although the nose provides about 90% of respiratory system air-
conditioning requirements [37, 38], further humidification and heating of the inhaled air occurs as it
travels through the pharynx and trachea. Fluid lining in these regions provides heat and moisture until the
air becomes fully saturated at core body temperature at a position termed as the isothermic saturation
boundary (ISB) [18]. The thermal and humidity gradient experienced by the inhaled/exhaled air, along
with the ISB location, fluctuate over the breathing cycle. Changes in ambient air temperature and
humidity also cause changes in ISB location.
AUTONOMIC REGULATION

Regulation and control of any process requires sensing. The thermo-receptors, concentrated around the vestibule, are believed to be the main sensors of inhaled air temperature. However, non-specific nasal innervation provides the ability to achieve strong tissue reactions, such as the sneezing reflex, under both normal and pathological conditions [39, 40].

Modulation of the nasal vascular flow by the autonomic nervous system regulates heat and fluid supply to the glands [41, 42] and adjusts airflow regime to meet air-conditioning demands [22, 43]. The ability of a healthy nose to condition extremes of environmental air to near alveolar conditions requires control of the heat and water fluxes across nasal mucosa. Achieving this requires simultaneous regulation of control of air flow regimes regulating heat and water mass transfer to and from the air, mucosal heat and water supply to airway fluid, as well as effective airway fluid transportation. These parameters are autonomically regulated through sympathetic and parasympathetic innervation [42], along with non-adrenergic, non-cholinergic gas signaling [44]. The control induced by this gas signaling process differs from that produced by the sympathetic and parasympathetic nervous innervation in a sense that it utilizes nitric oxide (NO) as a neurotransmitter [42, 44, 45]. NO is produced throughout the respiratory tract by the epithelial mucosa [33, 46-48]. Different isoforms of NO synthase occur within the respiratory tract under normal and pathological conditions [44]. For example, inducible nitric oxide synthase (iNOS) may assist microbiological defenses whilst endothelial nitric oxide synthase (eNOS) assists vasodilatation. Under normal conditions, nasal nitric oxide (n-NO) production is regulated through the absorption and synthesis of air-borne O2 by the nasal epithelial cells [49]. The output of n-NO is proportional to airflow [46, 50]. From an air-conditioning perspective, vasodilators (such as n-NO [47, 51]) or vasoconstrictors (such as antihistamine [52, 53]), either absorbed from the inhaled air or circulated within the blood, can influence blood flow within the nasal vasculature [54].

Airflow Regime

Within each nasal passage, the regulation of turbinate volume controls the airflow cross-sectional area (CSA), causing variation in the air velocity and flow regime that regulates heat and water mass flux. Commencing at the anterior region, during rapid inhalation, the CSA of the vestibule or “external nasal valve” is stabilized by cartilaginous tissue and inspiratory isometric contractions of the alar dilator
muscles which results in a relatively constant restriction to airflow [36]. The incoming airflow meets the
highest resistance when it reaches ‘the internal nasal valve’ region. Before entering the larger cavernous
nasal space, the incoming laminar flow accelerates as it passes through the reduced CSA, creating
turbulent airflow which enhances heat and water transfer within the nose [36, 55, 56].

Autonomic regulation of outgoing and recovered nasal fluid heat and water flux is realized through
changes in intranasal airflow resistance causing variation in air velocity distributions within each nasal
passage [57, 58]. This is achieved through independent regulation of blood flow through the cavernous
plexus or nasal erectile tissue found predominantly in the inferior turbinates and opposing septal wall [36,
56, 59]. This tissue consists of a network of large anastomosing veins capable of rapid blood volume
changes through the cushion or throttle activation in the regulating drainage veins [40]. Airflow
constriction occurs when the anterior end of the inferior turbinate projects into the nasal valve region,
causing a reduction in CSA [36]. The dynamic behavior of the internal nasal valve is demonstrated by its
erectile tissue reflex which can be initiated by applying pressure to one side of the body on a single armpit
or simply by lying on one side [60, 61]. Both of these situations result in blocking of the nasal passage on
the side on which the stimulation was applied. The functional purpose of this autonomic reflex, mediated
by skin pressure receptors, minimizes sweating on the downside of the body when lying on the side,
however, the regulation of the nasal erectile tissue appears altered as well. This reflex demonstrates that
regulation is at a central level, since other autonomic functions on the corresponding side are also affected
[62].

Part of nasal autonomic regulation involves ‘the nasal cycle’, which is a periodic congestion/decongestion
process of the erectile tissue alternating from one side of the nose to the other [63-65]. The cycle periods
range from 10 minutes to 8 hours and usually go unnoticed since the total nasal airflow resistance remains
unchanged [22, 42]. This variation in intranasal airflow patterns occurs in 20-40% of the population [58,
59] and coincides with an increase in fluid secretion in the patent side, but does not influence the degree
of water vapor saturation in the inhaled air [58, 63]. Currently it is thought that n-NO concentration
increases on the obstructed side, but in general the role n-NO plays in the nasal cycle remains uncertain
[66]. The duration of this ultradian cycle positively correlates with vasoconstrictor sympathetic outflow
and has been assessed by measuring the difference in brain hypothalamic and ear pinna temperatures in
cats [67]. In humans, electroencephalography (EEG) has demonstrated an alternating dominance in cerebral hemispheric activity that correlates directly with the nasal cycle [58, 62, 68]. The purpose of the nasal cycle is not fully understood; however, it may control the balance between heat and water fluxes from the airway fluid lining [69], as well as enable cells and glands on the constricted side to rest and recharge [42].

Mucosal Heat and Water Supply

A sub-epithelial network of fenestrated capillaries, positioned parallel to respiratory epithelium, supply both heat and humidification within the nose [40, 70]. At least half of the nasal blood flow passes through arteriovenous anastomoses (AVAs) which are connections between two blood vessels. These provide an alternative flow path during periods of congestion in erectile tissue. The role of AVAs is related to temperature and water control in the same way as in other parts of the body, such as in the hands and feet [70]. Mucosal blood flow is under autonomic nervous control with sympathetic nerves causing vasoconstriction and parasympathetic nerves dilatation [58]. The vasodilatory effect of n-NO contributes to the regulation of nasal blood flow by providing an additional means of controlling airway fluid secretion and heating [51, 71, 72].

The airway epithelium actively absorbs fluid, however, the estimated rate of submucosal gland secretion is significantly greater [73] which results in a net airway fluid gain. Submucosal gland mucus secretion and the depth of underlying PCL are regulated by autonomic mechanisms [40, 70, 74, 75]. Purinergic cellular fluid release and absorption also contribute to airway fluid volume regulation [26, 76]. Here fluid release occurs during multiple stress stimuli [77] and in response to airway fluid Na$^+$ and Cl$^-$ balance [24, 26].

Airway Fluid Transportation

During a breathing cycle, there is a continual flux of water and heat between the nasal airway fluid layer and inspired and expired air (Fig. 1). This fluid layer is a thermal and fluid buffer as well as a sink that traps inhaled particulate matter and water soluble toxins. The MTV normally ranges from 3 to 25 mm/min and it moves airway fluid toward the nasopharynx to ensure contaminant clearance and prevention of an excess mucus plaque build-up, which could obstruct water exchange between air and the PCL layer.
Changes in PCL rheology, tonicity, adenosine tri-phosphate (ATP) concentration, mucus depth and cell
temperature all influence ciliary beat frequency (CBF) [18, 29, 77-81], which in turn affects the MTV.
Recent research has found that n-NO also plays an important regulatory role in nasal air-conditioning
through regulation of CBF [47, 48, 82, 83], airway fluid transportation [71] as well as vascular derived
fluid and heat output [72]. CBF is, however, primarily regulated through purinergic pathways.

PURINERGIC REGULATION

Whilst most nasal mucus is produced by airway glands that respond to acetylcholine and vasoactive
intestinal peptide [84], it is also secreted from goblet cells. Airway epithelial cells respond, through
receptors and channels, to extracellular molecules and ionic concentrations that result in the blocking, or
secretion/absorption of specific fluids passing through the cell membrane. These receptors and channels
form the purinergic regulation system. Dynamic regulation of mucin secretion by goblet cells into the
mucus layer, release of intracellular ionic fluids that make up the PCL and cilial driving action are all
thought to be achieved through epithelial cell purinergic pathways [85]. As breathing stimulates
nucleotide activation and ion transport that achieves periciliary fluid volume regulation, the following
review is presented in this sequence.

Breathing Stimulation

Intracellular ATP release is triggered by mechanical deformation [86, 87], fluid shear stress [88, 89] and
compression or stretch of human airway epithelia [90, 91]. Extracellular release of this signaling molecule
occurs during oscillatory pressure fluctuations generated during normal tidal breathing. This stimulus is
important in the regulation of mucus clearance and the maintenance of a healthy airway [89, 91]. ATP
and other nucleotides from the airway epithelia are released by mechanical forces [26, 92] which are
impacted by breathing-induced cyclic shear and pressure stress which is exerted on the airway fluid layer
and oscillatory trans-epithelial pressure fluctuations [26]. Experimental work on tracheal epithelia has
shown that ATP release rates are most sensitive over the normal physiological range of oscillatory shear
and compressive stress [26]. While the airway wall shear stress during normal breathing is estimated as
high as 3 dyn/cm² in the nose [93] (compared to around 0.45 dyn/cm² in the trachea and lower airway
[89]), it is likely that the nasal epithelium responds to airway stresses in a similar fashion to the trachea.
Since goblet cell mucin and PCL volume is increased via epithelial ion transport and MTV enhanced by
an increase in CBF [85, 94-98], this stimulation is also considered part of a natural nasal defense system
to wash away noxious stimuli [77, 96].

Recent work on sheep trachea has found that fully humidified unidirectional airflow at 30°C or even 34°C
is insufficient to prevent epithelial cell dysfunction or damage from occurring [99]. During normal tidal
breathing healthy nasal epithelia is exposed to air well below these temperatures and humidity values.
This highlights the importance of cyclic air-stress stimulation in maintaining healthy ASL volume and
MTV.

**Nucleotide Activation and Ion Transport**

PCL volume regulation is achieved by activation of two classes of receptors, P1 and P2, in response to
extracellular stimulation by nucleotides and their metabolites [96], see Fig. 2. This activation opens
pathways to regulate specific ionic fluid transport across the epithelial cell wall. Currently, all four known
P1 receptors, A1, A2A, A2B, and A3, are primarily activated through adenosine [77, 85]. This metabolite is
derived from ATP that has been converted to adenosine mono-phosphate (AMP) and then adenosine via
hydrolysis by ecto-nucleotidases present on the cell surface [96, 100]. Activation of the A1 or A3
receptors releases intracellular K⁺ and Cl⁻ into the surrounding fluid. Variation in intracellular calcium
(Ca²⁺) occurs when a subgroup of P1 receptors, type 2Y, couple to a signal enzyme phospholipase C
(PLC), forming inositol 1,4,5-trisphosphate (IP₃). This acts as a secondary messenger molecule, releasing
Ca²⁺ from internal stores [94]. P1 receptor activation produces a discharge of intracellular fluid and an
elevated CBF [78, 101]. However, activation of A2A and A2B receptors releases intracellular Cl⁻ ions and
produces the same increase in intracellular IP₃ levels that releases Ca²⁺, producing elevated CBF.

Of the sixteen known P2 receptors, there are eight metabolic (P2Y) receptors, Y1, Y2, Y4, Y6, Y11, Y13 and
Y14. All are activated when extracellular concentrations of the nucleotides ATP, uridine 5'-triphosphate
(UTP) or their metabolites, ADP, UDP and UDP-sugars, reach sufficient concentrations [77, 102]. Like
the P1 receptors, activation of the P2Y group elevates intracellular IP₃ levels, which releases Ca²⁺,
elevating CBF, whilst simultaneously signaling the opening of channels that release intracellular K⁺ [103]
and Cl⁻ [104, 105] into the surrounding PCL. It is believed that a Ca²⁺ independent Cl⁻ channel exists [96]
and that the P2Y₂ and A₂B receptor groups have one of the strongest influences on CBF and airway ion
and water transportation [100, 101, 106-108]. Eight P2X receptors, triggered exclusively by ATP [77, 85], increase cell membrane Ca^{2+} and Na^{+} permeability [77, 104].

**Periciliary Liquid Volume Regulation**

Airway epithelia, characterized as having low transepithelial resistances [109] and high water permeability [110-112], finely tunes the PCL hydration state [113]. This is achieved through continual back-flux occurring during cellular ion and water release [26]. Leaky epithelia are capable of absorbing fluid through Na^{+} absorption and discharging fluid through Cl^{-} secretion. The PCL NaCl concentration is essentially isotonic with the epithelial cell [114]. It has been hypothesized [26, 115] that salt secreted by the epithelia is followed by water, to maintain tonicity, resulting in a PCL volume increase. Conversely, if salt is absorbed by the epithelia, water will follow into the cells, resulting in PCL dehydration. Data shows continuous switching of these ion channels between secretion and absorption phenotypes [89, 113]. Epithelial water flux possibly fine tunes PCL volume during tidal breathing.

**MODELING**

Heat and mass flow are essential for any air-conditioning process. Local heat and water mass transfer coefficients have been reported utilizing naphthalene sublimation techniques [116]. However, nowadays, computational techniques, such as computational fluid dynamics (CFD), are commonly used to predict these coefficients.

Simulation techniques of airflow, heat and water mass transfer are used to understand the influence of morphology and mucosal conditions on nasal air-conditioning and also to predict the influence of surgical interventions. Early ex-vivo physical models in cadaver half heads or casts visualized flow through transparent plates, which replaced the septum [117]. Although this technique was troubled by uncontrollable cadaver tissue shrinkage, it provided qualitative visualization of airflow patterns through smoke particles in air or dye filaments in liquid [22, 55, 117, 118]. Later the use of plastination overcame the problems associated with tissue shrinkage [119]. Quasi-static one-dimensional theoretical models developed to predict heat and water mass transfer [120] compare well with the temperature profile measured with in vivo studies [121]. Later two and then three dimensional computational studies utilizing simplified nose-like features enabled local determination of air temperature and humidity [122, 123]. On
the other hand, quantitative measurements within cadaver models using both laser Doppler velocimetry and miniature hot-wire anemometry enabled discrete air velocity measurement. Digital particle image velocimetry now suggest—that the 2-D approach does not provide an accurate representation of the complex 3-D airflows present within the complex nasal cavity [22].

Modern computer tomography (CT) and magnetic resonance imaging (MRI) techniques enable the capture of subject specific in vivo 3-D digital sectional nasal images [117, 124-126]. These techniques avoid the problem of cadaver tissue shrinkage encountered in ex-vivo physical testing and provide accurate complex nasal topographical data for use in computation fluid dynamic (CFD) models. Implementing four differential equations of conservation of mass, momentum, convection-diffusion and thermal energy balance into CFD modeling enables determination of instantaneous spatial distribution of air velocity, pressure, temperature and water vapor concentration along with heat and water flux [22, 122]. Mesh-based CFD models require sharp gradients in mesh density to maintain an anatomically correct model. This poses difficulties in mesh refinement if accurate results are to be obtained in these regions; however, the use of non-mesh based CFD overcomes this issue [117].

Assumptions of constant temperature and nasal wall water vapor concentrations commonly applied to CFD models ignore the spatial and time dependent distribution of the surface/gas interface and of the bulk fluid circulation [127]. Whilst some early CFD studies assumed steady airflow, during normal breathing the cyclic airflow causes variation in wall shear stresses and temperatures as well as in intranasal temperature [69, 128]. However, unsteady simulation utilizing 3-D CFD models overcomes this issue and supports previous estimates that the nose provides 90% of the heat and water flux required to modify inhaled air to alveolar conditions over an extreme range of ambient air temperature and humidity [122]. These results suggest that nasal air-conditioning is virtually independent of environmental conditions which is in support of previous in vitro studies [69]. On the other hand, pressure-flow measurements utilizing plastinated human facial specimens are comparable with those obtained by CFD techniques [119]. Ex-vivo plastinated and physical modeling along with CFD techniques have identified the nasal valve region as having the greatest influence on heat and water mass transfer due to induced higher airflow velocities [117, 119-121, 124-126, 128].
Recent CFD modeling of gas exchange between the maxillary sinus and nasal cavity predicted that for small single sinus ostia and large concentration gradients, diffusion was the dominant mechanism for sinus NO transport mechanism [129]; while for larger ostia and lower concentration gradients, convection mass transfer became dominant. Although modeling of n-NO content relative to nasal air pressure cannot be found in current literature, 3-D simulations utilizing a 6mm diameter ostium considering both diffusion alone and combined diffusion/convection produced results that matched testing of physical models [129].

An early clinical breathing therapy humidification model proposed that maintenance of optimum airway fluid rheology was necessary in order to achieve optimum mucociliary transportation [18]. This model, which includes tracheotomy breathing, assumed that inspired air temperature and humidity were the sole factors influencing airway fluid transportation and rheology. No mention was made of air pressure and this model in effect disregarded the air-conditioning contribution from the nose and trachea. Mathematical modeling of mucociliary clearance fluid mechanics that consider ciliary beat cycle, CBF, metachronous coordination and airway fluid rheology have been utilized to predict MTV [30]. Nasal wall distention under pressure has also been modeled [130] and was used to predict changes in airflow velocity during breathing at elevated air pressures. Numerous models utilizing fixed geometry of airway heat and water transfer have been undertaken [116, 120, 131-133], however, to the best of our knowledge, none considers the influence of many variables, such as changes in geometry and cell air stress-stimulation, due to breathing therapy air pressure on nasal air-conditioning.

AIR-CONDITIONING PATHOPHYSIOLOGY
The mechanisms through which nasal breathing therapy influences the efficacy of the nose to heat and humidify inhaled air are poorly understood. Pressurized breathing therapy, utilizing ambient air, frequently invokes negative nasal side effects. This suggests that the nasal mask conditions, in particular pressure, may be a contributing factor. However when reviewing patient symptoms, care needs to be given to subjective sensations. For example, perceived nasal patency may not be a reliable indicator of nasal congestion [134]; nasal inhalation of menthol triggers the sensation of improved nasal patency while airway resistance remains unchanged [59]. Compounding the difficulty in achieving reliable information is the buffering effect of the airway lining liquid, which produces a time delay between heat and water supply and loss/recovery to nasal air. A recent study has found no correlation between the
applied air pressure and the efficacy of nasal air-conditioning or related nasal symptoms during n-CPAP therapy [135]. However, testing was only undertaken over a 20-minute period and nasal airway liquid reserves probably made up for the shortfall in supply. An apparent contradiction to this suggestion is found in research into the long-term effects of n-CPAP therapy [28]. No changes in nasal resistance, mucociliary clearance or CBF was found before or after the application of long term breathing therapy. Researchers suggested that n-CPAP use might aggravate nasal inflammation, but unfortunately no data was collected during breathing therapy. In this study, the nasal mucosa may have recovered after cessation of treatment.

Mucosa Heat and Water Supply

The sub-epithelial network of fenestrated capillaries, positioned parallel to and facing the respiratory epithelium, are considered the humidification and heating source within the nasal cavity [40]. Nasal vascular beds making up the erectile tissue, glands and radiator vessels have a complex interlinking vasculature and nerve signaling structure [42]. This complexity makes it very difficult to identify discrete pathway or sensory regions responsible for changes in the mucosal heating and water state. Little information is available on the controlling mechanisms, both in terms of the affecting variables and sensors regulating heat and water transport within the nasal airway tissue layer. However, the vasodilatation effect of n-NO is thought to have a regulatory role in nasal air-conditioning [72, 136, 137].

Body position influences nasal air-conditioning with improved water availability and air heating sitting upright when compared to a supine position [138]. This is in contradiction with what researchers had predicted, namely, in the supine position, an increase in nasal blood volume in the venous sinuses occurs simultaneously with increased heat and water flux. The increase in nasal resistance, as indicated by a reduction in nasal volume due to swelling of nasal erectile tissue, does not appear as a consequence of increased nasal blood flow [43]. It seems that the capacitance vessels located deeper within the mucosa regulate nasal resistance and are independent of blood flow in the superficial mucosa layer which provides heat and moisture. This disassociation confirms earlier work on dogs, which concluded that nasal airway resistance cannot be correlated to vascular resistance or blood volume [139]. These results suggest that different parts of the mucosa respond differently to the same stimulus, such as changing body position. However the actual mechanism by which this occurs has not yet been established.
Perceived nasal dryness and the formation of mucus crusts may occur when water loss from the airway fluid exceeds its supply [40]. A possible cause is a reduction in the number or damage to the fenestrated capillaries within the nasal cavity. However, this does not explain why the use of nasal breathing therapy could cause these symptoms. On the other hand, vasomotor or non-allergic rhinitis, where nasal blood vessels dilate causing the nasal lining to fill with fluid and blood, is thought to occur as a consequence of excessive nasal fluid secretion [40, 140]. A change in the regulation of the musculature of these vessels is readily achieved through exposure to agents, such as vasodilators, circulating in the blood or by stimulation of the adventitial nerve endings in the outer blood vessel layer. Stimuli such as temporary humidity changes, ingestion of alcohol, infection, weather changes, stress, airborne irritants, medication and hormonal changes may all contribute to rhinitis [141-144]. Further, a reduction in plasma extravasation in airway epithelial cells, due to elevated air pressures, is believed to be due to a hydrostatic pressure-operated mechanism [145]. These findings in saline and histamine challenged mucosa may have no bearing on PCL supply in healthy mucosa during nasal breathing therapy.

A link exists between the ultradian nasal and sleep cycles which may influence nasal air-conditioning during breathing therapy [58, 62, 64, 68]. While awake and during synchronized or non-rapid eye movement (n-REM) sleep, vascular heat exchangers, including those found in the nasal mucosa, regulate body temperature through autonomic sympathetic vasoconstriction [67]. During desynchronized or rapid eye movement (REM) sleep, the pressure difference across the vessel walls tends to regulate blood flow with autonomic control becoming dysregulated, but not absent. During this sleep period, applied air pressure could reduce nasal vascular flow rates. This hypothesis is supported by the use of negative pressure to treat erectile dysfunction by facilitating an increase in blood flow to the penis [146, 147]. The effect elevated air pressure has on the nasal erectile tissue vasculature is not yet known.

**Airway-Fluid Transportation**

Efficient transport of the airway fluid is important in matching the heat and water flux gradients from the underlying cellular and glandular structures to cyclic airflow humidification requirements. Airway fluid transportation is provided by coordinated cilia beating at a frequency influenced by many factors including n-NO concentration [47, 48, 82, 83]. Allergic rhinitis increases n-NO levels but this is
decreased in chronic sinusitis [33]. An increase in airflow through the nose also increases n-NO release with higher levels occurring during inhalation than exhalation [50, 71], but due to lower airflow rates, lower release occurs during sleep [148]. Since n-NO production is modified by changes in nasal blood flow and nasal volume [48], this may well be in response to a greater air-conditioning demand being made on the nose during periods of high air flow. On the other hand, humming increases n-NO concentrations 5-15 fold when compared to silent breathing and this effect diminishes when repeated [47, 48, 149-151]. Physical variables such as the nasal cavity volume, posture and airflow resistance do not change n-NO levels whilst smoking, hypoxia, intense exercise and the use of NO synthase (NOS) inhibitors and decongestants cause a reduction [47]. It is believed that L-arginine supplementation is useful in increasing NO concentrations in some patients [83]. Hyperbaric research [152] has demonstrated a 16% reduction in n-NO levels during periods when breathing air elevated to a pressure of 49 cm H₂O. This is more than twice the maximum pressure normally experienced during CPAP therapy. Further, occlusion of the sinus from the nasal cavity could prevent sinus sourced NO from entering the nose space, leading to subnormal n-NO levels in some situations. Soft nasal erectile tissue has frequently been found in the regions adjacent to openings of the paranasal sinuses where the mucosa has formed a lip or margin [40].

CBF appears to be related to changes in epithelial cell size. In murine tracheal tissue, reduction in cell size and CBF correlates directly with changes in extracellular fluid tonicity [81]. Water loss reduces CBF whilst re-hydration restores this action. CBF decreases under hypertonic conditions and conversely increases after dilution to hypotonic levels. This occurs with a range of osmolytes, including sodium chloride, mannitol, xylitol, glucose, cesium chloride and ethanol. The demonstration of hypotonic stimulation is consistent with the finding that cell swelling stimuli causes extracellular ATP or UTP release [81, 96]; this is a known purinergic regulator of CBF [78]. Mechanical stimuli, possibly through elevated airway pressure forces, are also known to initiate this release [77, 96]. This could adversely influence the epithelial purinergic pathways regulating CBF and ion transport across the epithelial cell boundary. As previously mentioned; in healthy subjects the airway surface liquid is approximately isotonic, resulting in equal ionic concentration existing across cellular walls [114, 115]. Epithelial cell damage changes the ion content in the connective tissue of the airway walls [153], reinforcing the view
that cell stimuli or damage may induce intracellular ATP or UTP release, leading to dysfunction of
epithelial cell purinergic pathways.

Air pressures of up to 20cm H₂O are delivered during n-CPAP therapy [9] which is well above the normal
pressures of around -0.1 to +0.3 cm H₂O experienced across a healthy nose during normal oscillatory
breathing [117, 119, 154]. This significant pressure augmentation effectively exposes the epithelial cells
to a non-oscillatory stress that could interrupt normal ATP or UTP release [26], disrupting normal
purinergic regulation, and cause reduction of airway fluid levels and CBF.

An investigation into the morphological changes that occurred in the nasal mucosa of patients undergoing
n-CPAP therapy over a 3-10 month period has found that the epithelium in all patients underwent
fundamental changes [6]. These included epithelial cell shape changes as well as ciliary clumping and
conglutination. However, this was contradicted by a later investigation [28] which made the influence
breathing therapy has on the nasal mucosa morphology yet unclear.

Exposure to temperature extremes also impacts negatively on CBF; below 5°C, CBF virtually ceases
[155] while above 50°C, epithelial cell death occurs [156]. Exposure to cold dry air can lead to epithelial
shedding with symptoms of rhinitis, rhinorrhea and nasal congestion in people sensitive to temperature
stimuli [157]. This shedding occurs due to a relaxation response, induced by cooling, occurs in the nasal
mucosa [158]. These temperature extremes are not normally encountered during breathing therapy.

Nasal breathing when compared to oral breathing has an important role in maintaining upper airway
mucosal “wetness” [159, 160]. Airway fluid rheology plays an important role in mucosal transport [80].
One clinical model concluded that air inhaled during breathing therapy needed to be fully saturated at
core body temperature to maintain the mucociliary transport system in an optimum state [18]. This
recommendation was made for tracheotomy patients, where the main airway source of heat and moisture
had been by-passed, and also for CPAP users. For this situation, progressive reduction in airway humidity
causes fluid thickening, a slowing of MTV [29, 161, 162], a reduction in CBF and epithelial cell damage.
Under extreme conditions, low fluid levels within the lungs could lead to atelectasis and poor gas
exchange [18]. On the other hand, progressive over humidification causes thinning of the airway fluid and
a slowing of MTV due to a reduced cilia driving force. If prolonged, this leads to cessation of fluid
transport and the risk of fluid draining into the lungs [18]. In extreme cases epithelial cell thermal damage
could occur due to excess condensation. A prolonged exposure time outside the optimum range could
lead to more severe consequences.

Heat and Water Convection Coefficient
The rate of heat and water vapor transfer occurring between the nasal mucosa and respiratory air
throughout the breathing cycle is governed by the corresponding convection coefficient. Airflow velocity
has a significant influence on these coefficients and is dynamically regulated in each nasal passage
through engorgement of erectile tissue, located in the nasal valve [36].

Congestion of the turbinate mucosa increases air turbulence which serves to enhance heat and water mass
flux [59]. Additional heat from the engorged turbinates and surrounding erectile tissue was thought to
enable the inhaled air to achieve further moisture acquisition within the nasal cavity. Recent
simultaneous MRI, temperature and humidity measurements within the nose have contradicted these
earlier ideas by suggesting nasal patency is independent of air-conditioning within the normal levels of
turbinate volume change [163]. During the breathing of ambient air, variation in the middle and inferior
turbinate volumes was found not to have a significant influence on intranasal air temperature and
humidity levels. Although the range where changes in turbinate volume influences nasal air-conditioning
was not established, the study confirms the possibility of both purinergic and autonomic regulation
systems working together to regulate nasal air-conditioning. Even though the study did not examine the
effect of nasal air pressure, it showed that variation in turbinate volumes may play a greater role in
congestion than in heating and humidification.

As previously stated, the purpose of the nasal cycle is not fully understood but it likely controls the
balance between heat and water fluxes from the airway fluid lining [69], as well as enable cells and
glands on the constricted side to rest and recharge [42]. The cycle may be affected by several factors
including body position. Given a shift in position, from lying flat to sitting upright having less influence
on the nasal cycle than a shift from lying flat to lying on the side [60], changes in posture have a
significant role in determining the airflow through the nose by. Positional changes may also disrupt cell
and gland recuperation periods by forcing change in duration of the nasal cycle.

Changes in nasal airflow resistance between upright and supine positions have been attributed to
increased venous pressure reducing venous drainage from the erectile tissue located in the nasal valve
region [42, 43, 60]. Nasal venous pressure increases by 8 mmHg between the upright and supine
positions [59]. However, the air pressure used in breathing therapy can exceed this value and cause
compression of soft erectile tissue and may result in suppression of blood flow in these regions. In
support of this is the finding that a significant proportion of patients who undertake n-CPAP therapy
experience an increase in nasal valve area and volume[4]. Both of these parameters have a significant
influence on nasal airflow velocity and subsequent variation in local heat and water mass transfer
coefficients within the nose.

Vasomotor, non-allergic and allergic rhinitis are thought to cause nasal congestion due to erectile tissue
swelling [40, 59]. The influence this has on autonomic regulation is not well established. However, the
nasal congestion in most cases can spontaneously resolve through the application of a vasoconstrictor,
such as an α-agonist, indicating that the swelling is due to changing the vascular capacitance rather than
the accumulation of extra-vascular fluid [36]. The congested side of the nose tends to have a higher total
water content than the non-congested side [140].

The influence that the nasal blood flow can have on the nose is seen in postcoital rhinitis, commonly
termed ‘honeymoon rhinitis’, which occurs when sexual excitement elicits sneezing, rhinorrhea and nasal
obstruction immediately after sexual intercourse [164, 165]. Although hyperventilation may cause nasal
congestion, the exact cause of this phenomenon remains unknown. Engorgement of the nasal erectile
tissue may occur as part of a nervous response, which includes sexual arousal [166].

Nasal Cycle
Transmural pressure regulates vascular flow during desynchronized or REM sleep [67] and also reduces ultradian cycle duration. Elevated air pressure may influence nasal vascular flow causing disruption to nasal air-conditioning. Reduction in sleep quality could perhaps explain partly the high level of dissatisfaction and non-compliance with n-CPAP therapy. Nasal breathing is designed to alternate from side to side in an ultradian cycle. Forced unilateral nostril breathing can influence this cycle at a central level [167, 168]. Both the influence of the lack of alternating nasal flow on the dream cycle, which is necessary for quality sleep, and the effect of n-CPAP therapy has on brain wave patterns during sleep remain unknown.

FUTURE INVESTIGATIONS

In a healthy airway there are many parameters that serve to regulate the efficacy of the nose to condition inhaled air. These include, but are not limited to airway fluid transportation, airflow regulation, heat and fluid supply. Each serves to regulate the efficacy of the nose to condition inhaled air and is seemingly influenced in some way by nasal mask air pressure. By isolating and trending their respective pressure related response, a better understanding of the effect of air pressure can be established.

Airway fluid transportation

Decoupling the influences of n-NO and epithelial stress stimuli induced ATP release on CBF driving mucociliary transportation of the mucus and PCL is problematic. It is not known yet how CBF is influenced when acted upon simultaneously by both of these variables. Intranasal n-NO concentration is a function of airflow rate and it appears influenced by nasal cavity air pressure. Currently, n-NO concentrations as a function of breathing cycle phase [71], flow rate [46] or under hyperbaric condition [152] are known. However, an in vivo measurement of n-NO in vivo as a function of applied breathing therapy pressure over the breathing cycle is required.

The suggestion that CBF is suppressed by excessive pressure leading to an excessive cell stress stimulus [81], needs to be confirmed, along with the relationship, if any, between CBF and applied breathing therapy air pressure. This could be achieved through recording airway liquid levels and particle transport velocity during in vitro nasal tissue pressure stress stimuli. Further investigation is also required to
establish if changes occur in airway fluid tonicity as well as extracellular ATP and other nucleotides during breathing therapy.

Airflow Regulation

The heat and water mass transfer convection coefficients are strongly influenced by the airflow regime present within the anterior region of the nose. Airflow is regulated through activation of erectile tissue in the nasal valve region. Venous pressure used to achieve this activation has a range similar to that applied by some nasal breathing therapies [59], so air pressure may influence erectile tissue regulation. This could be investigated through the use of MRI to measure the \textit{in vivo} changes in nasal morphology that occur between breathing ambient and pressurized air.

Heat and Fluid Supply

Changes in nasal vasculature flow supplying mucosal heat and water could occur due to the vasodilator effect of n-NO as well as from the influence of air pressure acting on soft mucosa tissue creating venous flow constriction. Decoupling of these variables during \textit{in vivo} testing will be difficult, however, any influence n-NO and air pressure have on the nasal vasculature needs further investigation. Despite research finding reduced plasma extravasation occurring during elevated air pressures [145], further work is required to determine if this occurs in healthy mucosa during nasal applied breathing therapy. MRI perfusion techniques could be utilized to quantify pressure induced change in nasal vasculature flows.

Airway fluid tonicity appears to be linked to epithelial stress stimuli in the regulation of airway fluid layer height. Investigations using \textit{in vitro} nasal tissue exposed to pressure stress stimuli and simultaneous tonicity measurement could improve our understanding of airway water supply.

Clinical Implications

Using nasal mask conditions to supplement any shortfall in nasal air-conditioning requires an understanding of the effect air pressure has on the regulatory systems. Investigations into nasal airway fluid transportation, airflow regulation and heat and fluid supply may lead to a therapy temperature/pressure/humidification algorithm that optimizes these parameters for a prescribed therapy pressure. Optimization could lead to a reduction in titration pressure and improved treatment compliance.
Until further work is undertaken to improve our knowledge in this area, the requirement to deliver fully saturated air heated to core body temperature remains the safest treatment option.

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Figure 1. Schematic representation of human respiratory air-conditioning elements.
Figure 2. Breathing stress induced stimulation of extracellular nucleotide and metabolites which serve to regulate airway fluid volume and CBF.
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