EVALUATION OF UPPER AIRWAY (UA) ANTHROPOMETRY USING MAGNETIC RESONANCE IMAGING (MRI) AND LATERAL CEPHALOMETRY IN PATIENTS OF OBSTRUCTIVE SLEEP APNOEA (OSA) IN NORTH INDIAN POPULATION

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Aim To study the anthropometric measurements of UA in patients of OSA and assess the relation between UA anatomy and severity of OSA.

Introduction Anatomical and nonanatomical factors are implicated in OSA in which repetitive collapse of UA occurs. Scarcity of data regarding UA anthropometry in OSA exist especially in Indian population.

Materials and Methods A prospective observational case control study was carried in which 60 OSA (mild:n=20, moderate: n=20, severe:n=20) and control group (n=20) underwent MRI and lateral cephalometry of UA. Linear Regression analysis (univariate and multivariate linear regression) of various parameters was done to find out the factors correlated with OSA.

Results In MRI, distance between hyoid bone and posterior nasal spine (H-PNS) and hyoid and posterior pharynx wall near vertebral column (H-COL) was found to be statistically significant (mild=60.23 mm, moderate=68.72 mm, severe=77.26 mm: control=60.23 mm:p=0.001) and (mild=14.8 mm, mod=15.2 mm, severe=19.09 mm: control=11.5 mm:p<0.001). Laterolateral dimension of tongue and lateral pharyngeal wall thickness were found to be statistically significant and increases with severity of OSA. Significant narrowing was found at level 1 i.e., rhinopharynx level and low retropalatal oropharynx level i.e., level 3(p<0.001). In lateral cephalometry, distance between hyoid bone and mandibular plane increase with severity of OSA and was found to be statistically significant among all groups(mild=16.68 mm, mod=25.25 mm, severe=28.1 mm:control=16.15 mm:p-value<0.001). On univariate analysis, Epworth sleepiness scale (ESS), neck circumference, Modified Mallampati Score, anteroposterior and laterolateral dimension at level 3 and 4, hyoid bone position and laterolateral length of tongue were associated with severity of OSA. On multivariate analysis, the following parameters remained significant- anteroposterior and...
laterolateral dimensions at level 3, hyoid bone position, ESS and neck circumference.

Conclusion In OSA, significant alteration of anthropometry of upper airway occurs and is associated with severity of OSA.

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**THE EFFECTS OF SUPPLEMENTAL OXYGEN ON BLOOD PRESSURE IN OBSTRUCTIVE SLEEP APNOEA DURING CPAP WITHDRAWAL**

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**Introduction** Intermittent hypoxia and/or intermittent arousals are thought to be the two key pathological mechanisms in the development of hypertension in obstructive sleep apnoea (OSA). We aimed to investigate the effect of abolishing the hypoxia on the rise in blood pressure (BP) that has been shown to follow continuous positive airway pressure (CPAP) withdrawal in patients with OSA. In addition, we explored the effect of supplemental oxygen on obstructive events during sleep.

**Methods** Patients with OSA, established on CPAP ≥ 1 year, and with ≥ 4 hours/night usage, underwent a week of screening and an ODI ≥ 20 on 1 of 3 nights off CPAP. Patients then received overnight supplemental oxygen or air (via real or sham concentrators) at a flow rate of 5 l/min during 2 weeks off CPAP. After at least two weeks ‘washout’ back on CPAP, subjects crossed over. Treatment order was randomised.

The primary outcome was early morning home BP which was recorded daily in triplicate, and averaged over the penultimate three mornings prior to each patient visit. Patients underwent overnight home respiratory sleep studies (including nasal airflow) on night 14 of each treatment arm. Differences between treatment arms were analysed using paired t-tests or Wilcoxon rank tests as appropriate.

**Results** Twenty-five patients completed the study. Their mean ± standard deviation age was 63±7 years, mean BMI was 35.3±6.7 kg/m², median (interquartile range) ODI at diagnosis was 48/hour (25, 68), and 21 (84%) were male. Table 1 shows the Results of the primary outcome and the overnight sleep studies.

**Discussion** Supplemental oxygen abolished the rise in early morning blood pressure during CPAP withdrawal when compared to supplemental air. As expected, supplemental oxygen substantially attenuated intermittent hypoxia and had only a small non-significant effect on the apnoea hypopnoea index. Thus intermittent hypoxia appears to be the dominant determinant of the rise in morning blood pressure seen in patients with OSA, rather than any other consequence of the obstructive events.

**Abstract P218 Table 1** The results of early morning blood pressure and overnight sleep studies on supplemental oxygen and supplemental air. Paired t-tests were used for home early morning blood pressure and heart rate measurements. Wilcoxon rank tests were used for overnight home sleep studies.

<table>
<thead>
<tr>
<th>Supplemental oxygen</th>
<th>Mean difference in change in BP (95% confidence interval)</th>
<th>p</th>
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</thead>
<tbody>
<tr>
<td>Baseline</td>
<td>Follow-up</td>
<td>Baseline</td>
</tr>
<tr>
<td>systolic BP (mmHg)</td>
<td>129.2±14.1</td>
<td>136.1±14.9</td>
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<tr>
<td>Mean</td>
<td>78.3±6.5</td>
<td>84.3±7.6</td>
</tr>
<tr>
<td>Mean heart rate (bpm)</td>
<td>81.7±8.3</td>
<td>69.4±8.9</td>
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**THE USE OF ORAL MODAFINIL IN CHRONIC OBSTRUCTIVE PULMONARY DISEASE PATIENTS WITH CHRONIC HYPERCAPNIC RESPIRATORY FAILURE**

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Hypercapnic respiratory failure (HRF) is common in advanced COPD. The standard treatment is usually nasal ventilation (NIV) to reduce the blood carbon dioxide level acutely or regular nocturnal use in chronic cases. Not all patients tolerate such treatment and failures occur. Published studies are conflicting showing varying benefit and a 2013 Cochrane review concluded that there was no evidence of significant benefit in any of the measured parameters. Other studies suggest a reduction in re-admission and death at 1 year between regular nocturnal NIV use and standard care.

We have used oral modafinil 200 mg/day as a respiratory stimulant for chronic HRF in COPD without NIV. We present the data from the first 11 cases (6 out of study and 7 in our current open randomised crossover study). The study patients had documented HRF for 6–12 months with PaCO₂ ≥ 6.5 before entry but refused NIV. We present the data at baseline and day 10 and 40 of modafinil, including mean arterial oxygen and carbon dioxide, Daytime and overnight oxygen saturations along with spirometry. We compare these Results with 2 published studies of NIV reported, one after 4 weeks and the other 12 months.

The study showed a mean improvement by day 40 of +1.8 kPa in PaO₂ and a reduction in PaCO₂ by 2.7 kPa. Daytime saturations, improved by +15% and overnight oxygen saturations by +15%.