
Anshul Tripathi, PACE Junior Science College, Mumbai
Raj Ramnani, Jayshree Periwal International School, Jaipur

Abstract:

In a novel approach to diagnose Obstructive Sleep Apnea, electronic components, such as an Arduino Mega, a Bluetooth Transceiver, an accelerometer, and an air-aularity sensor, were put together to create a wearable that would detect the frequency of apnea events, detect and diagnose the disorder, and sound an alarm when necessary. A primary consideration was to make the mechanism accessible and affordable, and in doing so, lower the cases of Obstructive Sleep Apnea that go undiagnosed due to the cost and inconvenience associated with the traditional diagnosis method—a Polysomnogram. Bluetooth capability was an additional consideration so that the device would transmit data directly to an android smartphone, eliminating the need for an additional output mechanism. The total cost of the device, quite surprisingly, did not exceed $30, and therein rendered the device an accessible, affordable mechanism for diagnosis. Tests of the device on diagnosed patients yielded data consistent with the diagnosis, with a few false positives as a result of the excessive sensitivity of the sensors.
Introduction

Imagine suddenly waking up at night with a jolt, gasping for breath, surprised by the fact that you feel as if someone was choking you. This is something that people suffering with a serious case of sleep apnea have no choice but to become used to.

Sleep apnea is a chronic sleep disorder which results in frequent pauses in breathing/shallow breathing during the sleep of a person. Obstructive sleep apnea is a type of sleep apnea caused when the throat muscles occasionally relax and constrict the airway, therefore preventing breathing. In other words, tissues in the upper airways approach and nearly come into contact with each other, blocking the inward flow of air. Symptoms of this sleep disorder can be general weakness during the daytime, headaches, snoring, etc. The most common methods of diagnosis include polysomnogram (PSG) at sleep laboratories (overnight sleep studies) and sleep monitoring devices. Both of these courses of detection are very expensive and often inconvenient, and consequently, unobtainable by the financially weaker section of society.

The primary objective of this project was to create a device that has the ability to do the following:

1. It can diagnose a person with sleep apnea
2. It can detect the number and frequency of apnea events during a sleep session, and consequently, can calculate the Apnea-Hypopnea Index (an indicator of the severity of the case of apnea in a given patient, defined as the apnea events per hour of sleep)
3. It has an alarm system that can alert people in case an apnea attack lasts too long.
4. It can investigate the effects of the following on a person’s apnea attacks:
   a. Air Temperature
   b. Humidity
   c. Carbon Dioxide Concentration
5. It should be compatible with an Android phone, and therefore, have Bluetooth functionality

The aim of this project was to end up with an open source device that can be modified/enhanced and used for research purposes, as well as be used by a common man suffering from sleep apnea to diagnose the presence and severity of his condition. It is worth noting that the device in no way aims to replace the use of a breathing aid, such as a Continuous Positive Airway Pressure (CPAP) device, used by patients already diagnosed with the disorder; rather we aim to make the diagnosis possible so that someone suffering from Sleep Apnea can seek immediate medical assistance.
An understanding of the mechanism of the pathophysiology of Obstructive Sleep Apnea was deemed integral to the creation of the device and is therefore detailed below:

- Air enters the body through either the mouth or the nose and makes its way to the pharynx. The path thus far is termed the upper airway.

- Ordinarily, the back of the pharynx is not robust and would therefore collapse inward as a person inhales. However, dilator muscles in place prevent this from happening and keep the airway open.

- Under abnormal circumstances entailing one or more of the following:
  - Dysfunction of the upper airway dilator muscles
  - Disproportionate Pharyngeal Wall Compliance
  - Intrapharyngeal and Peripharyngeal Impediments

An Apnea event can occur, temporarily hindering breathing. Most events do not wake the person but some may cause them to intermittently gasp/snort for breath.

- Obstructive Hypopnea can be defined as incomplete obstruction of the upper airway, which leads to shallow breathing. This leads to snoring indistinguishable from that not caused by apnea, rendering diagnosis laborious.

- The continuation of the blockage of the upper airway entails a positive feedback loop wherein the dearth of oxygen in the blood signals the diaphragm to contract and therein draw air into the lungs. As inhalation increases, the blockage is exacerbated and can lead to dire consequences unless the dilator muscles regain control.

Parts Used

The following parts have been used for the creation of this device:

1. Arduino Mega ($9.50)
2. HC05 Bluetooth Transceiver ($4.50)
3. MPU6050 (3-axis gyroscope + 3-axis accelerometer) ($3)
4. 5 Volt Piezo Buzzer ($0.15)
5. DHT11 (Temperature + Humidity sensor) ($1.50)
6. MQ-135 Air Quality Sensor (One of the gases for which it can measure concentration is Carbon-Dioxide) ($4)
7. DS 1307 Real Time Clock (RTC) Module ($1.50)
8. Capacitive Touch Sensor ($2.50)
9. SD Card Module ($2)
All of these together have a cost under $30/Rs.2000

Functions and Working

- **Sleep Apnea Diagnosis/ Apnea Attack Detection**

  a. For the purpose of detection and diagnosis of sleep apnea, an accelerometer is used to measure chest movement. When the device is strapped around the chest, the accelerometer is in a position parallel to the chest plate.

  The accelerometer measures the force acting on it in 3 directions; the x-axis (along one edge of the accelerometer), the y-axis (along the edge perpendicular to the x-axis edge), and the z-axis (perpendicular to the plane of the accelerometer). When stationary and with its plane parallel to the surface of the earth (i.e. the person is lying flat on his back), the only force acting on the accelerometer is downward gravity ($F_G$). $F_G$ is also along the z-axis and therefore, $F_{Gz} = F_G$. When a person inhales, his chest exerts an upward force ($F_{Bz}$) perpendicular to the z-axis of the accelerometer. In the case that the person is not lying flat on his back, the accelerometer is at an incline. Since the accelerometer is always strapped to the chest, breathing always exerts a force along the accelerometer’s z-axis. However, gravitational force vector acts downwards and is broken into components $F_{Gx}$, $F_{Gy}$, and $F_{Gz}$. Therefore, in this case, the net gravitational force is calculated using the vector resolution formula

\[
F_G = \sqrt{[(F_{Gx})^2 + (F_{Gy})^2 + (F_{Gz})^2]}
\]

[Diagram of accelerometer plane parallel to Earth’s Surface and at an incline with Earth’s Surface]
The algorithm that detects breathing is able to do so with the help of this formula:

\[
F_z = \sqrt{[(FG_x)^2 + (FG_y)^2 + (FG_z)^2]} - F_{Bz}
\]

What is the meaning of this formula? Well, when stationary, \(F_{Bz} = 0\). However, during inhalation, when the chest exerts an upward force, \(F_{Bz}\) becomes greater than 0, and the value of \(F_z\) dips. This dip refers to one breath. For a person without sleep apnea, these dips are periodic. However, if the accelerometer goes for a relatively longer period of time without the detection of such a dip, then it can be inferred that the person has stopped breathing and is suffering from an apnea attack. Such a period of invariance can be called an invariance event.

It is worth noting that hypopnea events (incomplete OSA, where the airway is only partially obstructed) will not trigger a dip in \(F_z\), so the device would be incapable of detecting and diagnosing hypopnea.

**Self-Learning Algorithm** – This algorithm is executed during the first run of the device and establishes the average frequency of inhalation. This algorithm allows for versatility by allowing it to be customized to different individuals with different breathing patterns.

**Noise-Cancellation Algorithm** – The accelerometer, when programmed, shows the values of force in terms of gravitational force along each axis. This means that when stationary and parallel to Earth, \(F_{Gz} \approx 1.00\) G, \(F_{Gx} \approx 0.00\) G, and \(F_{Gy} \approx 0.00\) G, where the numbers mean 1 G force, 0 G force, and 0 G force respectively.

However, since this value is only precise to 2 decimal places and can practically vary only between -1.00 G to 1.00 G (it is impossible for the chest to generate a force greater than gravity, i.e. 1.00 G), it is not very accurate.
The dips in the wavy line above, which represent breathing, although visible, are very small in magnitude. Therefore, the preciseness needed to be increased.

Registers are the memory locations where the accelerometer values are stored. Upon some research, we find out that the raw values being pulled from the registers are being divided by a scaling factor of 16384. Therefore, we can remove this division operation and receive values ranging from -16384 to +16384. Therefore, we obtain a precision that is just over 163 times greater than when the raw values are divided by the scaling factor (it can even detect changes in its values if a feather fell on it).

However, owing to the variation and imperfection in the raw values, a great amount of noise is encountered.

This noise is visible in the above graph. Such a great amount of noise makes pattern recognition and dip detection difficult. Therefore, to counteract this and smooth out the noise to a manageable level, a noise reduction algorithm is used. This algorithm is as follows:

Let \( R_i \) be the \( i \)th reading of the accelerometer. Let \( R_{in} \) be the new data point in the set of noise-reduced values

Using this simple noise reduction algorithm and recursively running it through the data points about 3, the noise is relatively lesser, and patterns more noticeable.
This is how pattern detection becomes easier and more precise.

- **Calculation of Apnea-Hypopnea Index**

  b. The Apnea-Hypopnea index is used to quantify the severity of sleep apnea. It is defined as the total number of apnea events divided by the total number of hours of sleep.

  In this device, this is accomplished with the help of a touch sensor and a real time clock (RTC) module. When the user touches the touch sensor once, an interrupt is triggered and the program starts running. A counter variable keeps track of the number of invariance events (a period of time during which dips in the graph are not detected). Each invariance event points to an apnea attack. After the user has finished sleeping and wakes up, he touches the sensor again. The total number of apnea events is then divided by the difference between the start time and end time. The result number gives us the AHI.

  An AHI of >30 would signal excessive risk of acute myocardial infarction or a stroke and would alert the user accordingly.

  A limitation of the method used is the failure to calculate a person’s Respiratory Disturbance Index (RDI). The Respiratory Disturbance Index (RDI) indicates any form of breathing irregularities and could possibly account for hypopnea-triggered deviations. As a result, one’s AHI is always equal to or lower than one’s RDI. Since the RDI isn’t calculated, risk posed by irregularities not in the form of described dip would go unidentified.
• **Alarm System**

c. If left unattended and allowed to continue for extended periods of time, apnea events have the potential to lead to a heart attack, a stroke, or even cardiac arrhythmia.

To prevent such a mishap from occurring, an alarm system has been incorporated into this device. If an invariance event (and therefore an apnea attack) lasts for longer than 45 seconds, the brain might get oxygen deprived and due to this possibility, an alarm system is triggered. A 5V buzzer sounds an alarm, as well as the phone (which is connected to the device via Bluetooth).

• **Investigation of Surrounding Factors**

d. This device allows for the investigation of the effect of 3 different atmospheric characteristics on the device:

   i. **Temperature** – Using DHT11 Sensor
   
   ii. **Humidity** – Using DHT11 sensor
   
   iii. **CO₂ Concentration** – Using MQ-135 sensor

Each of the 3 elements being varied are measured in a room, with 3 different subjects suffering from Obstructive Sleep Apnea over a course of 7 days. The data is recorded on to an SD card and analyzed later. The results of the experiment are documented in a later section of this report.

• **Android Application**

e. A rudimentary android app is designed that serves the following purposes:

   i. Returns the hours of sleep, number of apnea attacks, and Apnea-Hypopnea Index.
   
   ii. Allows for the comparison of AHI on different days under different environmental conditions to assess the effect of said conditions on severity of Apnea.
   
   iii. Sounds an alarm when necessary

Given below are a few screenshots of the application:
Conclusion

3 test subjects known to be suffering from obstructive sleep apnea were taken for testing purposes. The device successfully detected apnea attacks at night. A few false alarms were detected regarding the unusually long apnea events as the device hung up and needed to reset. As for the effects of atmospheric elements, the results are as follows:

- **Temperature (controlled through AC & monitored through sensor)** – The average AHI for the 3 candidates over 2 days was 27 at 18 °C and 23 at 22 °C. IE as temperature increases, AHI decreases.

- **Humidity (controlled through humidity controller and monitored through sensor)** – No significant variation.

- **CO₂ Concentration (controlled through the number of people sleeping in a given room and monitored using sensor)** – The average AHI for the 3 candidates over 2 days was 23 at 510 ppm and 25.3 at 560 ppm. IE as CO₂ concentration increases, AHI increases.
### Temperature results

<table>
<thead>
<tr>
<th>Subject</th>
<th>Day 1 AHI (Control) (25°C)</th>
<th>Day 2 AHI (22°C)</th>
<th>Day 3 AHI (18°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subject 1</td>
<td>21</td>
<td>23</td>
<td>28</td>
</tr>
<tr>
<td>Subject 2</td>
<td>20</td>
<td>21</td>
<td>23</td>
</tr>
<tr>
<td>Subject 3</td>
<td>24</td>
<td>25</td>
<td>30</td>
</tr>
</tbody>
</table>

### Humidity results

<table>
<thead>
<tr>
<th>Subject</th>
<th>Day 1 AHI (Control) (70%)</th>
<th>Day 4 AHI (40%)</th>
<th>Day 5 AHI (80%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subject 1</td>
<td>21</td>
<td>21</td>
<td>22</td>
</tr>
<tr>
<td>Subject 2</td>
<td>20</td>
<td>21</td>
<td>20</td>
</tr>
<tr>
<td>Subject 3</td>
<td>24</td>
<td>24</td>
<td>24</td>
</tr>
</tbody>
</table>

### CO₂ Concentration Results

<table>
<thead>
<tr>
<th>Subject</th>
<th>Day 1 AHI (Control) (440 ppm)</th>
<th>Day 6 AHI (2 people in room) (510 ppm)</th>
<th>Day 7 AHI (3 people in room) (560 ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subject 1</td>
<td>21</td>
<td>23</td>
<td>27</td>
</tr>
<tr>
<td>Subject 2</td>
<td>20</td>
<td>21</td>
<td>22</td>
</tr>
<tr>
<td>Subject 3</td>
<td>24</td>
<td>25</td>
<td>27</td>
</tr>
</tbody>
</table>

### Bibliography:


