Model – Based Approach to Estimate dFRC in the ICU Using Measured Lung Dynamics

A.N. Mishra*, Y.S. Chiew*, J.G. Chase*, G.M. Shaw**

*University of Canterbury, Christchurch, 8041, New Zealand (e-mail: ankit.mishra@pg.canterbury.ac.nz, yeong.chiew@pg.canterbury.ac.nz, geoff.chase@canterbury.ac.nz)

**Christchurch Hospital, Christchurch, 8011, New Zealand

Abstract: Acute Respiratory Distress Syndrome (ARDS) is characterized by inflammation and filling of the lung with fluid. Mechanical ventilation (MV) is used to treat ARDS/ALI using positive end expiratory pressure (PEEP) to recruit and retain lung units, thus increasing pulmonary volume and dynamic functional residual capacity (dFRC) at the end of expiration. However, simple methods to measure dFRC at the bedside currently do not exist and other methods are too invasive and impractical to carry out on a regular basis.

Stress-strain theory is used to estimate ∆dFRC, which represents the extra pulmonary volume due to PEEP, utilizing readily available patient data from a single breath. The model uses commonly controlled or measured parameters (lung compliance, plateau airway pressure, PV data) to identify a parameter \( \beta_1 \) as a function of PEEP and tidal volume. A median \( \beta_1 \) value is calculated for each PEEP level over a cohort and is hypothesised as a constant throughout the population for the particular PEEP. Estimated ∆dFRC values are then compared to measured values to assess accuracy of the model.

∆dFRC was calculated for 9 patients and compared to the measured values. The median percentage error was 40.29% (IQR = 14.20, 55.39) for PEEP = 5 cmH2O, 31.12% (IQR = 10.53, 192.71) for PEEP = 10 cmH2O, 20.8% (IQR = 7.51, 81.06) for PEEP = 15 cmH2O, 15.44% (IQR = 11.92, 36.18) for PEEP = 20 cmH2O, 19.7% (IQR = 4.79, 20.76) for PEEP = 25 cmH2O and 11.78% (IQR = 2.99, 27.5) for PEEP = 30 cmH2O. Linear regression between estimated and measured ∆dFRC produced \( R^2 = 0.862 \).

The model-based approach offers a simple and non-invasive method which does not require interruption of MV to estimate dFRC. The clinical accuracy of the model is limited but was able to track the impact of changes in PEEP and tidal volume on dFRC, on a breath-by-breath basis for each PEEP.

Keywords: Functional Residual Capacity, dFRC, FRC, Pulmonary, Model, ARDS, Intensive Care, ICU

1 INTRODUCTION

Acute Respiratory Distress Syndrome/Acute Lung Injury (ARDS/ALI) are characterized by inflammation and fluid filled lungs. Severely affected lung units collapse and cannot be recruited without external intervention, resulting in a smaller, stiffer lung, called the “baby lung” (Gattinoni and Pesenti, 2005). Mortality rates for ARDS have been reported between 30% - 70% (Bersten et al., 2002, Luhr et al., 1999, Manzano et al., 2005, Suchyta et al., 1997).

Clinicians offer a supportive environment that aids patient recovery by application of mechanical ventilation (MV), which partially or completely takes over the patients breathing effort. The primary focus of MV is to improve recruitment of lung units and increase gas exchange, while minimizing further harm (Ware and Matthay, 2000), primarily through application of a suitable positive end expiratory pressure (PEEP).

PEEP is important since ARDS affected lung units are vulnerable to collapse due to the extra pressure of the fluid and denaturing of surfactant that maintain the shape of the alveoli. PEEP prevents alveolar collapse and enhances gas exchange using higher pressure to recruit the collapsed alveoli. However, high PEEP and tidal volume (\( V_t \)) can also damage healthy alveoli (Bersten, 1998).

Functional Residual Capacity (FRC) is the lung volume at zero end expiratory pressure (ZEEP). PEEP ensures pulmonary volume above FRC. This additional volume, dynamic FRC (dFRC) is schematically shown in Figure 1:

Fig. 1. dFRC at PEEP > 0 and FRC at ZEEP
dFRC offers useful clinical information regarding the lung recruitability, but is not normally measured at the bedside. Some mechanical ventilators allow FRC and dFRC measurements (GE Healthcare., 2006), but most do not. However, these measurements require interruption of MV and can be additionally invasive. Computer Tomography (CT) scans timed at the end of expiration allow accurate assessment of FRC (Malbouisson et al., 2001). However, it has limited application since it requires transport of the patient out of the Intensive care unit (ICU), disconnecting the patient from the ventilator and exposure to radiation. Other methods include washout of tracer gases, such as Nitrogen (Heinze et al., 2007, Olegard et al., 2005), which allows precise measurement of FRC, but requires the use of a dispensing device for the tracer gas, negating its frequent use at the bedside (Weismann et al., 2006).

dFRC is clinically important, can indicate lung condition, and, if tracked regularly, can identify changes in patient condition. The model proposed in this research aims to estimate dFRC from readily available pressure-volume (PV) data without interrupting MV treatment. The model is an extension to a previous model proposed by Sundaresan et al (Sundaresan et al., 2011) which required PV data at a minimum of two PEEP levels. More PV loops at a wide range of PEEP levels were required for higher accuracy. The new model proposed here requires PV data from only one breath at one PEEP level to estimate dFRC at different PEEP levels for a given patient. This major extension allows continuous tracking of dFRC at the patient’s bedside without interrupting MV.

2 METHODS

2.1 Model Summary

Chiumello et al (Chiumello et al., 2008) proposed a stress-strain theory of lung dynamics which was used by Sundaresan et al (Sundaresan et al., 2011) to develop a model to estimate dFRC. Chiumello et al defined transpulmonary pressure (ΔP_L) as the clinical equivalent of stress. Transpulmonary pressure is the difference between the applied airway pressure and the corresponding pleural pressure. The clinical equivalent of strain is the ratio of the change in volume (ΔV) to the FRC, which represents the resting lung volume, yielding a stress-strain definition:

\[ ΔP_L = E_{Lspec} \times \frac{ΔV}{FRC} \]  

Where, the specific lung elastance (E_{Lspec}) can be defined as the transpulmonary pressure at which FRC effectively doubles. The values of specific lung elastance reported by Chiumello et al (Chiumello et al., 2008) were 13.4±3.4 cmH2O for the control subjects, 12.6±3.0 cmH2O for the medical control group, 14.4±3.6 cmH2O for the ALI subgroup, and 13.6±4.1 cmH2O for the ARDS subgroup. The study indicated that specific lung elastance does not vary significantly across subgroups.

Transpulmonary pressure can also be defined in terms of plateau airway pressure ΔP_{aw}:

\[ ΔP_L = ΔP_{aw} \times α \]  

Where:

\[ α = \frac{E_L}{E_L + E_{CW}} \]  

Where α in Equation 3 represents the static lung elastance, defined by the Lung Elastance (E_L) and the Chest Wall Elastance (E_{CW}). Chest wall elastance plays an important role in MV as part of the airway pressure applied is used to inflate the lungs and the rest is utilized to overcome chest wall pressure. This relation and the effect of chest wall elastance is shown in Figure 2.

It can be observed that the total elastance is the same in both cases in Figure 2. Case (a) is typical of ARDS patients where a stiffer lung is represented by the higher lung elastance compared to case (b), which shows a healthy lung. Hence, the value of α indicates the severity of ALI or ARDS, where a higher value of α indicates a higher severity of ARDS.

![Fig. 2. Effects of different lung and chest wall elastance (Gattinoni et al., 2004)](image)

The values of α reported by Chiumello et al (Chiumello et al., 2008) were 0.69±0.15 for control group patients, 0.74±0.16 for intensive care patients with medical diseases, 0.64±0.15 for the ALI subgroup and 0.71±0.16 for ARDS subgroup.

Combining Equations 1 and 2 yields:

\[ FRC = \frac{ΔV}{ΔP_{aw}} \times \frac{E_{Lspec}}{α} \]  

Equation 4 defines FRC as a function of the lung compliance (ΔV/ΔP_{aw}), E_{Lspec} and α of the patient.

It was hypothesised that dFRC follows similar mathematical form as Equation 5. It should be noted that ΔdFRC in Equation 6 represents the additional volume in the lung above FRC due to application of PEEP and not the absolute dFRC, which consists of the FRC as well.

\[ FRC + ΔdFRC = \frac{ΔV}{ΔP_{aw}} \times \frac{E_{Lspec}}{α} (1 + x) \]  

Defining dFRC = FRC + ΔdFRC yields:

\[ ΔdFRC = \frac{ΔV}{ΔP_{aw}} \times \frac{E_{Lspec}}{α} x \]  

Where x is a function of the PEEP level at which ΔdFRC is estimated. E_{Lspec} and α are constant parameters and can be combined into one parameter, β, yielding:
\[ \Delta \text{dFRC} = \frac{V_t}{\Delta \text{Paw}} \times \beta \]  

(7)

Where \( \Delta V \) is replaced by tidal volume (\( V_t \)) and \( \beta \) is a function of the PEEP applied, \( E_{\text{lspec}} \) and \( \alpha \). The assumption that \( \alpha \) is constant is true only for the linear portion of the PV loop (Sundaresan et al., 2011).

Sundaresan et al (Sundaresan et al., 2011) proposed a model to estimate the \( \Delta \text{dFRC} \) defined:

\[ \text{dFRC} = \frac{\Delta \text{dFRC}}{\Delta \text{PEEP}} \times \beta \]  

(8)

Where \( \frac{\Delta \text{dFRC}}{\Delta \text{PEEP}} \) represents the volume responsiveness of the lung to a change in PEEP and \( \Delta \text{dFRC} \) represents the additional volume due to PEEP. Sundaresan et al (Sundaresan et al., 2011) also hypothesized that the value of \( \beta \) for a particular level of PEEP can be assumed as a population constant for a given PEEP, which is used here.

2.1.2 Patients

Clinical data for 9 patients (Sunderesan, 2010) in Table 1. The data is characterized by different levels of lung injury, including PV data for at least 3 PEEP levels per patient. The data did not contain information on the FRC of the lung but contained the measured \( \Delta \text{dFRC} \) values for each PEEP level for each patient. This study uses an average breath for each PEEP level. The use of data was approved by the New Zealand South Island Regional Ethics Committee.

2.1.3 Analyses & New Approach

The proposed model estimates \( \Delta \text{dFRC} \) based on the compliance (\( V_t/\Delta \text{Paw} \)) observed in the PV data recorded for each patient. A representative breath, with PEEP level and corresponding peak inspiratory pressures normally observed in clinical settings was selected to calculate the lung compliance for each patient since the lung compliance was found to change with pressure. In particular, it was found to decrease at higher PEEP levels. Since the FRC values were not measured, \( \frac{E_{\text{lspec}}}{\alpha} \) could not be estimated. Instead, the \( \beta \) was calculated using Equation 8. Calculated \( \beta \) values were then normalized by tidal volume (\( V_t \)) as dFRC can vary with the \( V_t \) applied, yielding:

\[ \beta_1 = \frac{\beta}{V_t} \]  

(9)

A median \( \beta_1 \) value was calculated for each PEEP level over all patients, which was used as a population constant. The \( \Delta \text{dFRC} \) value for each PEEP level for each patient was then estimated by substituting this median \( \beta_1 \) value into Equation 7, and multiplying the resultant value with the applied \( V_t \). The median and (IQR) of errors are reported for all (\( N = 53 \)) data points (patients and PEEP levels).

The linear trend observed in median \( \beta_1 \) values at each PEEP level over the cohort used to linearly interpolate median \( \beta_1 \) values at intermediate PEEP levels to generalise the approach. These interpolated \( \beta_1 \) values were then used to re-estimate \( \Delta \text{dFRC} \) values at all the PEEP levels, using only surrounding data to interpolate from.

Overall, this new approach and analysis can be summarised:

- Calculate \( \beta_1 \) for each patient and PEEP
- Use median \( \beta_1 \) for each PEEP as a population constant to estimate \( \Delta \text{dFRC} \) at those PEEP values for each patient
- Assess error between calculated and measured \( \Delta \text{dFRC} \) values.

This approach requires values for \( \beta_1 \) from patient data at all PEEP values that might be used. To generalise this approach, interpolating \( \beta_1 \) between known \( \beta_1 \) values at different PEEP levels was tested. Thus:

- Using median cohort \( \beta_1 \) values for two surrounding or near PEEP values, an interpolated \( \beta_1 \) is calculated for each PEEP level.
- Interpolated \( \beta_1 \) for each PEEP as a population constant to estimate \( \Delta \text{dFRC} \) at those PEEP values for each patient
- Assess error between calculated and measured \( \Delta \text{dFRC} \) values.

These results test a more general approach. Performance is assessed by trend correlation coefficient (R), where.

This approach is unique from the work of Sundaresan et al (2011) in that this prior work required multiple PEEP values to estimate dFRC, which was not practical clinically.

3 RESULTS

Table 2 shows the analytical solution for Median [IQR] \( \beta_1 \) for each patient and PEEP and overall cohort results. Figure 3 shows the variation of median \( \beta_1 \) values with respect to PEEP. Figure 4 shows the trend between estimated and measured \( \Delta \text{dFRC} \) values with a \( R^2 = 0.862 \) using this population constant. Figure 5 shows the general trend observed between the re-estimation and measured \( \Delta \text{dFRC} \) using interpolated \( \beta_1 \) values with \( R^2 = 0.850 \). Finally, Table 3 shows the percentage error (Median [IQR]) between measured and estimated \( \Delta \text{dFRC} \) values.

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<tr>
<th>Sex</th>
<th>Age [years]</th>
<th>Cause of lung injury</th>
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<tbody>
<tr>
<td>Patient 1</td>
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</tr>
<tr>
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<td>Male</td>
<td>22</td>
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<tr>
<td>Patient 3</td>
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<td>Patient 5</td>
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<td>Patient 6</td>
<td>Male</td>
<td>69</td>
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<td>Patient 7</td>
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<td>56</td>
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<td>Patient 8</td>
<td>Female</td>
<td>45</td>
</tr>
<tr>
<td>Patient 9</td>
<td>Male</td>
<td>61</td>
</tr>
</tbody>
</table>

Table 1: Characteristics of the patient studied
Table 2: β₁ values calculated for each patient based on the representative breath chosen, where 4_2 and 6_2 are second trial on the same patient 3 and 8 days later respectively

<table>
<thead>
<tr>
<th>PEEP [cmH₂O]</th>
<th>5</th>
<th>10</th>
<th>15</th>
<th>20</th>
<th>25</th>
<th>30</th>
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<td></td>
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<td>1</td>
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<td>0.0160</td>
<td>0.0317</td>
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<td>2</td>
<td>0.0083</td>
<td>0.0209</td>
<td>0.0338</td>
<td>0.0480</td>
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<td>3</td>
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<td>0.0385</td>
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</tr>
<tr>
<td>4</td>
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<td>0.0070</td>
<td>0.0214</td>
<td>0.0352</td>
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<td>5</td>
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<td>0.0307</td>
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<td>0.0059</td>
<td>0.0190</td>
<td>0.0355</td>
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<td>8</td>
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<td>0.0233</td>
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<td>9</td>
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<td>0.0076</td>
<td>0.0195</td>
<td>0.0322</td>
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<td>Median</td>
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<td>0.0328</td>
<td>0.0480</td>
<td>0.0545</td>
<td>0.0606</td>
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<tr>
<td>IQR</td>
<td>0.0057, 0.0105</td>
<td>0.0072, 0.0235</td>
<td>0.0200, 0.0396</td>
<td>0.0352, 0.0546</td>
<td>0.0483, 0.0678</td>
<td>0.0558, 0.0821</td>
</tr>
</tbody>
</table>

Fig. 3. Median β₁ vs PEEP. The median β₁ here is calculated based on the lung compliance of a representative PV loop.

Fig. 4. Estimated vs measured ∆dFRC values using exact β₁ values from the cohort at each PEEP.

Fig. 5. Measured vs Estimated ∆dFRC values using interpolated β₁ values.

4 DISCUSSION

The model was developed to estimate ∆dFRC as a function of PEEP and tidal volume using only readily available PV data and to do so at a single PEEP setting to avoid interrupting MV treatment. It estimates ∆dFRC using the compliance observed by applying the stress-strain approach proposed by Chiumello et al (Chiumello et al., 2008). Overall, dFRC offers important information on the status of the lung and its recruitability, and using such a non-invasive approach it can be tracked continuously as it changes with the evolution of disease using the proposed method.

It was also observed in Figure 3 and Table 2 that median β₁ values change almost linearly with respect to PEEP at each PEEP level. This linear trend allows the linear interpolation of median β₁ values at intermediate PEEP levels. Thus, to generalise this approach a b value is not needed at every possible PEEP. There was almost no loss in fidelity or correlation as observed comparing the R² values for the results in Figures 4 and 5.
Table 3: Percentage error between the observed and measured dFRC values

<table>
<thead>
<tr>
<th>PEEP [cmH₂O]</th>
<th>5</th>
<th>10</th>
<th>15</th>
<th>20</th>
<th>25</th>
<th>30</th>
</tr>
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<td>Patient</td>
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<td></td>
<td></td>
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<tr>
<td>1</td>
<td>44.18</td>
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<tr>
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<td>0.94</td>
<td>0.00</td>
<td>6.87</td>
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<tr>
<td>3</td>
<td>0.93</td>
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<td>6.17</td>
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<tr>
<td>4</td>
<td>527.41</td>
<td>198.63</td>
<td>68.62</td>
<td>36.18</td>
<td>23.22</td>
<td>11.78</td>
</tr>
<tr>
<td>5</td>
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<td>311.93</td>
<td>177.03</td>
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<tr>
<td>IQR</td>
<td>14.20</td>
<td>55.39</td>
<td>10.53</td>
<td>192.71</td>
<td>7.51</td>
<td>81.06</td>
</tr>
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</table>

Sundaresan et al (Sundaresan et al., 2011) reported \( R^2 = 0.946 \) over the entire cohort. However, a significant drawback in the model proposed by Sundaresan et al was that it could not be used for continuously tracking dFRC since it requires PV data for at least 2 PEEP levels. Its application in real-time dFRC measurement is thus limited without interrupting MV treatment and consuming clinical time.

The new model proposed here utilises readily available clinical data from a single breath. It can thus be automated and does not interrupt MV therapy nor require any special clinical intervention. These advantages allow continuous tracking of \( \Delta \text{dFRC} \) and thus of dFRC if FRC is measured. The compromise in accuracy is acceptable to maintain this advantage.

However, it must be noted that the method proposed has some limitations in its predictive capability. In some cases, the percentage error observed between the estimated and measured \( \Delta \text{dFRC} \) values was high, up to almost 500% in one of the patients studied. Larger errors were primarily found at very low PEEP levels, below Auto – PEEP (Sunderesan, 2010). Such low levels are generally not observed in clinical settings. Equally, Auto-PEEP can be detected directly from PV loop responses. Thus, this issue is manageable. In particular, ignoring low PEEP values less than known Auto-PEEP levels (Sundaresan et al 2011b) yields maximum errors of only XXXX%.

In general, larger estimation errors can limit the use of this or any similar method for estimating the recruitment potential. However, median percentage errors in all cases were 10-30% and generally lower as PEEP rose above Auto-PEEP. Equally, the trends were still valid.

In particular, estimated \( \Delta \text{dFRC} \) follows a similar trend with respect to PEEP as the actual \( \Delta \text{dFRC} \). Hence in spite of higher error values in limited cases, the model is still capable of predicting the trend in \( \Delta \text{dFRC} \) with respect to PEEP. This particular outcome is shown in Figure 6, which shows the measured and estimated \( \Delta \text{dFRC} \) values for Patient 5 where percentage errors observed were high at low PEEP values due to an Auto-PEEP of XXX cmH₂O. It is clear that while there

![Fig. 6. Measured and Estimated dFRC for patient 5](image_url)

is bias, the trend is quite accurate, which would allow effective clinical assessment and use.

In conclusion, the model presented allows real-time assessment using data from only a single PEEP level. Thus, there needs to be no interruption to MV therapy or any added clinical effort to obtain this value – a completely non-invasive, model-based approach. The accuracy of the resulting values is only slightly reduced from methods that require several PEEP levels and significant, invasive clinical intervention. The trends over time and PEEP are equally accurate ensuring that there is no loss in clinical applicability and value.

5 CONCLUSIONS

A novel method of estimating dFRC for a given mechanically ventilated patient is presented, based on a stress-strain approach. The method presented allows real-time assessment using data from only a single PEEP level. Thus, there needs to be no interruption to MV therapy or any added clinical effort to obtain this value – a completely non-invasive, model-based approach. The accuracy of the resulting values is only slightly reduced from methods that require several PEEP levels and significant, invasive clinical intervention. The trends over time and PEEP are equally accurate ensuring that there is no loss in clinical applicability and value.
REFERENCES


