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The computation of reference state and APE production by diabatic processes in an idealized tropical cyclone

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This study investigates the energetics of tropical cyclone intensification using the Available Potential Energy (APE) theory. While the idea is now well accepted that tropical cyclones (TCs) intensify as the result of the conversion into kinetic energy of the APE generated by the release of latent heat extracted from the warm tropical ocean surface, its rigorous theoretical formalization has remained elusive owing to the complexity of constructing a suitable reference state for defining and quantifying APE in a moist atmosphere. Yet the construction of such a reference state is a key fundamental issue, because the magnitude of the APE reservoir and of its temporal evolution, as well as the values of the thermodynamic efficiencies controlling the rate at which diabatic processes generate or destroy APE, depend on its specification. This issue is illustrated in the idealized context of an axisymmetric TC model by comparing the energetics of TC intensification obtained by using two different sorting-based approaches to compute the reference state defining APE. It is found that the thermodynamic efficiency controlling the APE generation by surface latent heat fluxes is larger when the reference state is constructed using a ‘top-down’ sorting method, as the APE thus defined absorbs all the CAPE present in the system. However, because a large fraction of the overall CAPE is never released during the TC’s lifetime (e.g. in regions dominated by subsidence), there is a better agreement between the production of APE by surface fluxes and its subsequent conversion into kinetic energy when a ‘bottom-up’ reference state is used. These results suggest that, contrary to what is usually assumed, the reference state in APE theory should be constructed to minimize, rather than maximize, the total APE, so that the introduction of dynamically inert APE is minimized.

Key Words: tropical cyclone; available potential energy; reference state; CAPE; intensification

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1. Introduction

A mature tropical cyclone (TC) is often characterised by a well-developed secondary circulation. At the surface, air flows cyclonically towards the vortex core. The surface convergence is balanced by rising motion in deep convection within the eyewall, followed by anticyclonic outflow in the upper troposphere and subsidence in regions at larger radii. As surface air spirals inward, sensible and latent heat are extracted from the ocean surface through surface fluxes. The latent heat is later released as the air rises within the eyewall which strengthens the convective updraughts, leading to stronger surface inflow and eventually maintains a thermally direct secondary circulation (Emanuel, 1991, 2003, give detailed reviews).

Emanuel (1986) first suggested that the secondary circulation of a TC vortex can be viewed as a four-stroke Carnot engine that converts energy acquired from the ocean into mechanical energy. In brief, as a surface air parcel moves down the pressure gradient towards the vortex core, it extracts energy from the ocean surface and expands isothermally. Upon reaching the base of the eyewall, the air parcel rises within the deep convection where it expands and cools adiabatically. The air parcel then enters the outflow channel near the tropopause. As it moves away from the vortex core, it compresses isothermally and loses energy to space by radiative cooling. The circulation is finally completed by sinking back to the surface at large radius where the parcel is warmed by adiabatic compression. In this view, the maximum amount of work that can be performed is given by the Carnot efficiency defined as $(T_s - T_{out})/T_s$, where $T_s$ represents the parcel’s temperature in the surface inflow and $T_{out}$ is its temperature in the upper-tropospheric outflow (Ozawa and Shimokawa, 2015, give a recent extension of this idea).
From an energetic viewpoint, the Carnot approach can be used to simplify the vortex’s energy budget into a two-term balance between energy production and mechanical dissipation. For example, Emanuel (1988) proposed the ‘Maximum Potential Intensity’ (MPI) theory that aims to predict the maximum possible intensity of a TC in a given atmospheric state. The MPI theory assumes that all the energy input is converted into mechanical energy at the Carnot efficiency, which is in theory the highest efficiency possible. Therefore, by equating such energy conversion to the mechanical dissipation one can predict the maximum possible intensity of the vortex. Smith et al. (2008) give a critical review of the MPI theory.

Another interesting characteristic of Emanuel’s Carnot approach is the realization that surface energy fluxes can be linked to their conversion into mechanical energy by the thermodynamic efficiency. The thermodynamic efficiency in Emanuel’s Carnot approach is determined by the temperature of the surface inflow and that in the upper-tropospheric outflow (i.e. $T_s$ and $T_{out}$). In Emanuel’s original framework, air parcels are assumed to circulate in a fixed path such that $T_s$ and $T_{out}$ are assumed to be constant. Generally, $T_s$ is mostly controlled by the sea surface temperature while locally it can be affected by localized processes such as precipitation downdraughts. Meanwhile, $T_{out}$ is mostly determined by how high an air parcel can rise in the eyewall convection, or simply the buoyancy of an air parcel. However, in a more realistic setting when air parcels do not follow the same trajectory in the secondary circulation, the thermodynamic efficiency is likely to vary across the vortex depending on the buoyancy of individual air parcel.

As well known from classical thermodynamic theory, the Carnot theory of heat engines arises from the consideration of the entropy budget applied to a system undergoing a closed thermodynamic cycle. As a result, a TC can be regarded as a Carnot heat engine only when viewed over its entire life cycle for the assumption of a closed cycle to be approximately satisfied. The approach is therefore difficult to apply to the study of the transient evolution and intensification of TCs. The theory of Available Potential Energy (APE), first introduced by Lorenz (1955) for understanding how the large-scale atmospheric circulation is maintained against dissipation, represents in principle a more satisfactory approach to study the energetics of stratified fluids (e.g. Tailleux, 2013a). Physically, this is because APE is by definition the fraction of the total potential energy convertible into kinetic energy (KE), and hence ultimately responsible for the observed intensification. Historically, APE has been primarily defined as the difference between the potential energy of the actual state minus that of a notional reference state obtained by means of a (moist) adiabatic rearrangement of the fluid parcels minimizing the total potential energy of the system. In the case of a dry and statically stable atmosphere, Lorenz (1955) demonstrated that the APE resides only in the horizontal gradient of temperature and pressure such that the reference state can be represented by the horizontal average of pressure and temperature fields.

For a moist atmosphere, Lorenz (1978, 1979) and Randall and Wang (1992) have argued that the presence of boundary-layer parcels with Convective Available Potential Energy (CAPE) complicates the definition of APE, as it makes it possible for the atmosphere to possess APE even for a horizontally homogeneous barotropic state. This implies, therefore, that CAPE induces a vertical component to APE. The connection between the two concepts was discussed in much detail in Emanuel (1994). However, since it is not generally possible for all the CAPE that exists at any given time in the atmosphere to be released, it seems obvious that absorbing all the CAPE into the definition of moist APE, as proposed by Lorenz (1978, 1979) and Randall and Wang (1992) may result in a value of APE possibly greatly overestimating the amount of potential energy that can be converted into KE. Physically, the APE theory would have a greater predictive value if APE could be defined to represent the ‘right’ fraction of the total potential energy actually convertible into KE, which is the basic idea explored in this article.

Lorenz (1978) suggested that one way to construct the reference state for a moist atmosphere is to use a ‘sorting’ procedure that rearranges all air parcels in the atmosphere diabatically into a hydrostatic and statically stable column. Various studies such as Randall and Wang (1992) and Tailleux and Grandpeix (2004) used similar diabatic rearrangement to compute the reference state and APE of a conditionally unstable atmospheric column, but it is unclear how such techniques can be applied to a realistic three-dimensional atmosphere. Also, note that the APE theory and the concept of reference state here are defined globally over the entire volume of the atmosphere. Meanwhile, CAPE is usually defined locally for a given air parcel by lifting it along a moist adiabat under the action of a positive buoyancy anomaly with respect to the local sounding without altering the surrounding atmosphere. Therefore, although APE may include contribution from CAPE in a moist atmosphere, they are two completely different concepts. Recently, the possibility of using a more general reference state in conjunction with a locally defined ‘APE density’ has been discussed by Tailleux (2013a,2013b) and Peng et al. (2015), but the APE density will not be covered in this study.

As far as we are aware, Pauluis (2007) is the only study which fully generalizes the discussion of sinks and sources of APE for a general moist atmosphere, but it provides no details about how to construct the reference state on which the theory relies. Assuming that a suitable reference state can be constructed, the discussion of energetics in APE theory takes the form

$$\frac{d\text{APE}}{dt} = G - C_{\text{APE\rightarrow KE}},$$

$$\frac{d\text{KE}}{dt} = C_{\text{APE\rightarrow KE}} - D,$$

where $d/dt$ is the total time derivative, $\text{APE}$ and $\text{KE}$ represent volume-integrated values of APE and KE, and $D$ denotes dissipation by viscous processes. Equation (1) shows that the change in APE in the atmosphere (dAPE/dt) is controlled by the balance between the production of APE by diabatic processes such as surface fluxes ($G$) and the conversion of APE into KE ($C_{\text{APE\rightarrow KE}}$). Pauluis (2007) also presented a detailed investigation of the production of APE by diabatic processes in a moist atmosphere. In general, the production of APE is governed by the addition (or removal) of heat and moisture by diabatic processes multiplied by a thermodynamic efficiency defined as $(T_i - T_{ref})/T_i$. This is very similar to Emanuel’s Carnot efficiency with $T_i$ and $T_{ref}$ replaced by the in situ temperature of an air parcel ($T_i$) and its temperature in the reference state after the adiabatic rearrangement ($T_{ref}$). Note that, for a dense air parcel that is rearranged downward in the reference state, $T_{ref}$ can be higher than $T_i$, resulting a negative thermodynamic efficiency. Another key difference is that each air parcel in the atmosphere will have a different position in the reference state, leading to a range of thermodynamic efficiencies depending on their buoyancy. For the sake of clarity, the thermodynamic efficiency $(T_i - T_{ref})/T_i$ will be referred to as the ‘APE production efficiency’ in this study. KE, on the other hand, grows as a result of the conversion of APE into KE, and decays as the result of viscous dissipation $D$, usually assumed to be primarily controlled by surface drag.

Based on the APE theory, this article aims to investigate the production of APE by diabatic processes and its conversion into KE in numerical simulation of an idealised TC vortex. The
complexity associated with finding a suitable reference state of a moist atmosphere with the adiabatic rearrangement approach is also addressed. The choice of the reference state is important, because it affects the values of the APE reservoir, as well as of the generation term \( G \) in Eq. (1). In contrast, the conversion of APE into KE (\( C_{APE\rightarrow KE} \)) is independent from the choice of reference state. To understand why it matters, let us integrate Eqs (1) and (2) over the life cycle of the TC, assuming the KE to be small at the beginning and end of the life cycle. This yields

\[
\Delta APE = G - C_{APE\rightarrow KE},
\]

\[
0 \approx C_{APE\rightarrow KE} = D,
\]

where \( \Delta APE \) represents the net change of APE over the life cycle of the TC and \( G, C_{APE\rightarrow KE} \) and \( D \) are the production, conversion and dissipation terms integrated over the volume and time. Combining these two equations yields

\[
G \approx D + \Delta APE.
\]

This formula is similar to that underlying MPI theory, but for the presence of the net APE change term \( \Delta APE \). Physically, one would expect \( \Delta APE \) to be negligible in order to be able to directly link generation and dissipation, but this is not necessarily the case depending on how the reference state is constructed. For instance, the initial and end states could both possess significant amounts of CAPE that is not released due to the presence of convective inhibition. Absorbing such inert CAPE in the definition of APE could give the illusion of a physically meaningless large value of \( \Delta APE \).

The main question investigated in this article is whether it is possible to construct the reference state such as to minimize \( \Delta APE \), which would establish the possibility of using APE theory to provide an alternative rigorous theoretical construction of MPI theory. Two different sorting methods with contrasting views on buoyancy are developed and used to analyse the output from an idealized TC numerical model. The production of APE by surface fluxes in the simulated TC is also investigated using the Pauluis (2007) APE framework. The production of APE by surface sensible and latent heat fluxes is computed and compared to the conversion into KE.

This article is divided into the following sections. Section 2 gives a brief introduction to the numerical TC model and the sorting methods. Section 3 shows the main results of the investigation followed by a conclusion and discussion in section 4.

## 2. Methodology

### 2.1. Axisymmetric tropical cyclone model

This study used a modified version of the axisymmetric TC model (hereafter 2D model) designed by Rotunno and Emanuel (1987). Modifications to the rain-physics scheme and gravity wave filter following Craig (1996) and Craig and Gray (1996) were also included. The simulation follows the typical ‘prototype problem’ set-up. The atmosphere was the mean hurricane season sounding (Jordan, 1958) and a weak initial vortex with maximum surface wind speed no stronger than 15 m s\(^{-1}\) was placed over an open ocean surface. The sea surface temperature was fixed at 29 °C throughout the simulation.

The 2D model contains nine prognostic variables including velocities in the radial, azimuthal and vertical directions (\( u, v, w \)), potential temperature (\( \theta \)), a dimensionless Exner pressure deviation from initial state (\( \pi \)), and the mixing ratios of water vapour (\( r_v \)), cloud water (\( r_c \)), rain water (\( r_i \)) and ice (\( r_i \)). The prognostic equations were integrated forward in time with a time step of 6 s for a simulation duration of 150 h. A shorter time step (0.6 s) was used for advancing \( w, \theta, \pi \) to account for acoustic waves. No parametrization scheme was used for convection as it is simulated explicitly. The 2D model domain extends radially to 3600 km and vertically to 27.5 km with a horizontal grid spacing of 2.5 km and vertical grid spacing of 0.625 km. Note that each grid box in the axisymmetric domain represents a ring-shaped volume around the centre of rotation. A no-flux boundary condition was used while ‘sponge layers’ were placed at the top and outer boundaries for the absorption of gravity waves. Surface fluxes were simulated using the bulk aerodynamic formulae. For sensible and latent heat flux, the coefficients were set at 0.001 and 0.0012. For surface friction, the drag coefficient (\( C_D \)) was computed using

\[
C_D = 1.1 \times 10^{-3} + 4 \times 10^{-5} \times (u^2 + v^2)^{1/2},
\]

in which \( u \) and \( v \) are evaluated at the lowest model level.

### 2.2. Behaviour of the simulated vortex

The evolution of maximum surface wind speed and minimum surface pressure of the simulated vortex can be divided into three distinct stages (Figure 1(a)). The first 40 h was a stagnation stage with little to no intensification. This was followed by a rapid intensification stage between 40 and 60 h. The maximum surface wind speed increased from just above 20 m s\(^{-1}\) at 40 h to nearly 70 m s\(^{-1}\) at 60 h while the minimum surface pressure

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Simulation results: (a) time series of minimum surface pressure (blue) and maximum surface wind speed (dashed red); (b) distribution of horizontal wind speed in the vortex after 120 h of simulation.
The definition of the reference state and how it can be derived from the observed atmospheric state is a vital part of the APE theory. As discussed by Lorenz (1955, 1978), one possible approach is the ‘sorting’ method in which air parcels in the atmosphere are rearranged adiabatically according to their buoyancy. Eventually, this will produce a hydrostatic and statically stable column of air such that no more total potential energy can be used to support it. As recently considered in the oceanic case by Saenz et al. (2015), there are two natural sorting strategies, namely the ‘top-down’ and ‘bottom-up’ sorting methods.

Consider a two-dimensional atmosphere with horizontal size $r$ containing four air parcels (Figure 3(a)). To construct the reference state, the top-down sorting method begins by lifting the observed atmospheric state to an equivalent potential temperature $\theta_e$ of the lowest model level. From 40 h onwards, the surface layer within 500 km radius is characterised by negative values of $u$, representing the surface inflow branch of the secondary circulation. However, there were also several fluctuations in radial velocity in the surface layer at 20–30 h (200–900 km radius), 60–80 h (200–2000 km radius) and 100–120 h (1500–2000 km radius) respectively. These fluctuations are marked by positive values of $u$, indicating a reversal in the surface flow direction. They tend to form at about 250 km radius and then propagate away from the centre of the vortex at a fairly constant speed of 25–35 km h$^{-1}$.

Comparing Figure 2(a) and (b) shows that the fluctuations in surface radial velocity clearly affected the values of $\theta_e$ in the surface layer. While the highest values of $\theta_e$ (>365 K in the mature vortex) can be found near the centre of rotation, the region beyond 500 km radius is dominated by continuous variation between higher (>355 K) and lower (<350 K) values of $\theta_e$ throughout the simulation. The variation is characterised by a gradual increase in $\theta_e$ values to a maximum of roughly 360 K. The time of maximum $\theta_e$ is usually observed during the onset of each surface radial velocity fluctuation. An abrupt drop in $\theta_e$ to below 350 K is then observed after each surface radial velocity fluctuation, followed by a gradual recharge until the next fluctuation. The clear outward propagation signal suggests that the fluctuations in $u$ and $\theta_e$ might be related to processes such as gravity wave or outward-moving rain bands associated with convective downdraughts, although more work would be required to confirm that such a propagation is not a numerical artifact.

Figure 2. Radius–time plot of (a) surface radial velocity ($u$) and (b) equivalent potential temperature ($\theta_e$) at the lowest model level.

Figure 3. Graphical illustration of the top-down and bottom-up sorting methods. See text for explanation.
The parcel with the lowest density after the relocation (parcel 1) is assigned to the pressure level \( P_{\text{ref,bot}} \) and removed from the pool of parcels. The same process is then repeated with parcels 2 to 4 at the next pressure level \( P_{\text{ref,bot}} + \delta P \) and then continuing downward (Figure 3(c)). Eventually each air parcel is assigned to a pressure level, resulting in a sorted air column that represents the reference state (Figure 3(d)). However, one could argue that the reversal of the top-down sorting method can also produce a valid reference state. In this case, the bottom-up sorting method constructs the reference state by bringing all four parcels downward to the surface represented by \( P_{\text{ref,bot}} \). The least buoyant air parcel (parcel 4) is assigned to this pressure level and removed from the pool. The process is then repeated with parcels 1 to 3 at the next pressure level \( P_{\text{ref,bot}} - \delta P \) and continuing upward until the construction is finished. Note that, although both sorting methods give the same result in Figure 3(d), this is not always the case in practice due to the presence of CAPE and each sorting method will produce a different reference state. This is the major focus of this article and will be demonstrated further in the next section.

Although the two sorting methods appear to be polar opposites, their operating principles are identical. In brief, grid boxes of the 2D model at a given integration time are regarded as individual air parcels. For both sorting methods, the values of \( \text{in situ} \) air temperature \( (T_i) \), pressure \( (P_i) \) and total mixing ratio \( (r_{\text{tot}}) \) are used to compute the liquid potential temperature \( \theta_i \) of all air parcels in the model domain. The air parcels are then relocated adiabatically and reversibly to the starting \( P_{\text{ref}} \) (e.g., \( P_{\text{ref,bot}} \) or \( P_{\text{ref,top}} \)) by conserving \( \theta_i \) and mass. The choice of using \( \theta_i \) as a conserved variable in the adiabatic relocation is because \( \theta_i \) is defined as an exact proxy for moist entropy \( (\eta) \) such that it satisfies \( \eta(\theta_i, P_i, r_{\text{tot}}) = \eta(T_i, P_i, r_{\text{tot}}) \), where \( P_i \) is the surface pressure. This coincidence with the definition presented in Emanuel (1994) if the air parcel brought adiabatically to \( P_i \) is unsaturated.

The parcel's temperature after the relocation, \( T_{\text{ref}} \), as well as the partition between vapour and liquid mixing ratio, \( r_v \) and \( r_l \), are then computed. An iteration procedure is described using the fact that \( \theta_i \) and \( r_{\text{tot}} \) are conserved in the relocation (i.e. \( \theta_i(T_i, P_i, r_{\text{tot}}) = \theta_i(T_{\text{ref}}, P_{\text{ref}}, r_{\text{tot}}) \)) such that \( T_{\text{ref}} \) and the new vapour–liquid partition can be computed iteratively from a range of possible values. The densities of the air parcels after the relocation \( \rho \) are then computed and the densest \( (\text{or lightest for the top-down sorting method}) \) parcel is selected and assigned to the current \( P_{\text{ref}} \) level. The next \( P_{\text{ref}} \) level is then computed using \( P_{\text{ref}} - \delta P \) \( (\text{or } P_{\text{ref}} + \delta P \text{ for the top-down sorting method}) \). The quantity \( \delta P \) represents the depth of the selected parcel after stretching it horizontally to match the radius of the model domain \( (\text{i.e. } r \text{ in Figure 3(c))} \) which can be calculated using the mass of the selected parcel and the hydrostatic relationship. After removing the selected air parcel from the pool, the remaining parcels are relocated to the updated \( P_{\text{ref}} \) thus starting the next relocation cycle. This process is repeated until all air parcels are assigned to a \( P_{\text{ref}} \) slot, resulting a hydrostatic and statically stable column of air parcel representing the reference state.

The pressure at the bottom and top of the reference state, \( P_{\text{ref,bot}} \) and \( P_{\text{ref,top}} \), can be expressed as \( P_{\text{ref,bot}} = P_{\text{ref,bot}} - \Delta P \). Both \( P_{\text{ref,bot}} \) and \( P_{\text{ref,top}} \) can be computed using the relationship between surface pressure and the total mass of air over a given surface area, while \( \Delta P \) can be computed using the total mass of atmosphere in the model domain and the hydrostatic relationship. Both terms can be expressed as

\[
P_{\text{ref,bot}} = \frac{\sum (P_i \times A_i)}{A_{\text{tot}}},
\]

\[
\Delta P = \frac{\sum (m_i \times g)}{A_{\text{tot}}},
\]

where \( P_i, A_i \) and \( m_i \) are the surface pressure, surface area and mass of the \( i \)-th column of the 2D model domain, \( g \) is gravity and \( A_{\text{tot}} \) is the total surface area of the model domain. As both \( P_{\text{ref,bot}} \) and \( P_{\text{ref,top}} \) (and any other \( P_{\text{ref}} \) levels in the sorted column) are horizontally homogeneous such that no more horizontal pressure gradient force (PGF) can further contribute to the APE in the atmosphere. Since mass is conserved in the sorting process, the top and bottom values of the reference states created by both methods are identical. However, the \( P_{\text{ref}} \) levels in between could be different if the sorting methods produce a different parcel permutation in the reference state.

Both the top-down and bottom-up sorting methods were used to analyse the output from the 2D model. To improve computational efficiency, only the lower portion of the model domain was analysed, while a large number of grid boxes representing the stratosphere were ignored. The cut-off height was set at 15 km above sea level such that the analysis includes most, if not all, of the troposphere and the important features of the simulated vortex.

### 3. Results

#### 3.1. Reference states

Figure 4 shows the reference state position of each air parcel produced using both sorting methods at 60 and 120 h into the simulation. The reference state position of an air parcel is represented by the pressure slot \( P_{\text{ref}} \) it occupies in the sorted column. For both sorting methods, a small \( P_{\text{ref}} \) value indicates an air parcel is relatively buoyant compared to the rest of the domain and is lifted upward in the reference state. Similarly, a large \( P_{\text{ref}} \) value suggests an air parcel is relatively dense and is relocated towards the surface in the reference state.

Using the top-down sorting method, buoyant air parcels with small \( P_{\text{ref}} \) values can be found near the surface at 60 h when the vortex has just finished its rapid intensification (Figure 4(a)). The surface parcels were able to reach a small \( P_{\text{ref}} \) because the CAPE stored in the boundary layer was released when the surface parcels were lifted upward during the top-down sorting. Also, parcels between 6 and 10 km above sea level near the centre of rotation were mostly lifted to smaller \( P_{\text{ref}} \) than those at larger radii, indicating this part of the vortex was relatively buoyant compared to the region further away from the centre of rotation. While surface parcels rely on the release of CAPE during the adiabatic rearrangement to reach a small \( P_{\text{ref}} \), parcels near the centre of rotation are in general thermally buoyant due to the warm core signature of the vortex and therefore lifted upward in the reference state.

At 120 h, the vortex has further developed and reached a near-steady state intensity. At this time, the surface layer is still very buoyant with most parcels lifted to small \( P_{\text{ref}} \), especially in the inner region within 1000 km radius. However, the number of parcels with small \( P_{\text{ref}} \) values has reduced slightly in the region between 1000 and 2000 km radius (Figure 4(b)). This could be related to a surface radial velocity fluctuation which reduced the \( \theta_i \) of the region at 120 h (Figure 2(b)).

At 8–14 km above sea level, the inner region of the vortex remained relatively buoyant compared to the rest of the model domain with the number of air parcels showing small \( P_{\text{ref}} \) values increasing slightly as the vortex intensified. Another notable difference compared to the earlier time is the presence of model columns within 1000 km radius which have smaller \( P_{\text{ref}} \) values than the surrounding environment. These buoyant air columns are rings of weaker convection outside the eyewall in which the release of latent heat increases the buoyancy of air parcels, resulting in a smaller \( P_{\text{ref}} \) in those model columns.

Similarly, the value of \( P_{\text{ref}} \) was computed using the bottom-up sorting method at both 60 and 120 h (Figure 4(c) and (d)). Above the boundary layer, the results at both integration times are generally similar to those produced by the top-down sorting method. The mid to upper-atmosphere (beyond 5 km above sea level...
level) within 1000 km radius is characterised by an anvil-shaped region of buoyant air parcels with $P_{\text{ref}}$ values smaller than 200 mb. Also, model columns with smaller $P_{\text{ref}}$ values can be found within 1000 km radius just outside the eyewall at 120 h which is similar to the top-down result described previously. However, the most remarkable difference from the results with the top-down sorting method when using the bottom-up method is the lack of buoyant parcels with small $P_{\text{ref}}$ values near the surface. At 60 h, only two localised patches of parcels with small $P_{\text{ref}}$ values are found in the surface layer at 800 and 1200 km radius. At 120 h, the number of surface parcels with small $P_{\text{ref}}$ values increased in the inner region, but the surface layer further away from the centre of rotation is still dominated by parcels with larger $P_{\text{ref}}$ values than in the top-down results.

To further investigate the different distributions of air parcels in the reference states computed by the two sorting methods, the model domain was divided into three regions representing the surface layer (i.e. lowest model level), the inner region (radius ≤ 1000 km) and the outer region (radius > 1000 km). Figure 5 shows a scatter plot of $P_{\text{ref}}$ against the in situ pressure ($P_1$) for both sorting methods at 120 h. Each grid box is coloured according to its origin using the division above.

In general, the distribution of air parcels in the reference states computed by the two sorting methods are very similar. Buoyant air parcels in the inner region (red) were lifted upward to $P_{\text{ref}}$ slots that are smaller than their in situ pressure while air parcels in the outer region (green) showed only small displacement as they are relatively stable. Similar to the difference in the values of surface $P_{\text{ref}}$, the most remarkable difference between the two sorting methods is the distribution of surface parcels (blue) in the reference state. A large number of surface parcels were lifted to the top of the reference state between 300 and 100 mb when using the top-down sorting method while most of them remained near the surface when using the bottom-up sorting method. Another notable difference is that, when using the top-down sorting method, some parcels from the surface and the inner region were relocated to the middle of the reference state between 600 and 400 mb; this was not observed when using the bottom-up sorting method.

As the buoyancy of the surface air parcels is largely determined by the amount of CAPE they possess, the different results for both sorting methods at 120 h.
produced by the two sorting methods are clearly related to the treatment of CAPE during the adiabatic relocation. In the top-down sorting method, the starting pressure \( P_{\text{ref, top}} \) is usually around 30 mb which is way above the typical lifting condensation levels (LCLs) of the surface parcels. Therefore, when the surface parcels were brought adiabatically upward to the upper troposphere, they were able to release most, if not all, of the latent heat they have. This has a huge impact on the density of a surface air parcel after the adiabatic relocation since the potential contribution from surface CAPE is included. Therefore, the top-down sorting method offers a maximizing view on the buoyancy of surface air parcels by assuming all CAPE in the surface layer can be released in the sorting process.

In contrast, when air parcels were brought to the surface in the bottom-up sorting method, they were warmed by adiabatic compression. As a result, air parcels were mostly unsaturated after the relocation and were thus unable to release latent heat. Therefore, only the most buoyant surface parcels were able to reach their LCL after the \( P_{\text{ref}} \) slots below the LCL were filled by denser surface parcels from the outer region beyond 1000 km radius. As a consequence, the bottom-up sorting method generally excludes the potential contribution from CAPE when evaluating the buoyancy of surface air parcels, leading to the lack of surface parcels with small \( P_{\text{ref}} \) values. Compared to the top-down sorting method, the bottom-up method offers a more minimizing view on the buoyancy of surface air parcels by placing a much harder restriction on the release of CAPE. Since both sorting methods were able to identify the buoyant parcels in the upper part of the inner region, it is clear that the construction of the reference state based on a parcel-sorting approach is not affected by the thermal buoyancy that is directly available, but the treatment of potential buoyancy associated with CAPE is more challenging. The results here highlight the complexity of finding the reference state of a moist atmosphere due to the presence of CAPE. In the next section it will be showed that the contrasting view on buoyancy from the sorting methods has a large impact on the computation of APE production by surface fluxes.

3.2. APE production efficiency

After constructing the reference state, the APE production efficiency, defined as \((T_1 - T_{\text{ref}}) / T_1\), is computed for both sorting methods. In general, the APE production efficiency reflects the buoyancy of an air parcel compared to the rest of the atmosphere since the difference between \(T_1\) and \(T_{\text{ref}}\) is determined by how the parcel is rearranged in the reference state. A buoyant air parcel that is relocated upward in the reference state will have a lower \(T_{\text{ref}}\) than the in situ temperature \(T_1\) leading to a positive APE production efficiency. In contrast, a dense air parcel that is relocated towards the surface will have a higher \(T_{\text{ref}}\) and thus a negative APE production efficiency.

Consistent with the distribution of parcels with small \( P_{\text{ref}} \) values, air parcels with the highest APE production efficiency were found in the surface layer and in the mid to upper-troposphere inside the inner region when using the top-down sorting method at both 60 and 120 h (Figure 6(a) and (b)). The APE production efficiency of individual surface parcels can be as high as 0.3, while most of the buoyant parcels in the mid to upper-troposphere is characterised by a weak negative APE efficiency. This is because the relatively stable air parcels in the outer region were relocated downward to make room for the buoyant air parcels from the surface and the inner region that were relocated to the top of the reference state. The compression and warming caused by the downward relocation resulted a larger \(T_{\text{ref}}\) than the in situ temperature \(T_1\), and thus gave a negative APE efficiency.

As the bottom-up sorting method restricts the release of surface CAPE during the sorting process, most of the surface parcels
smaller more surface parcels were relocated upward and attained a much higher average surface APE production efficiency since methods. In general, the top-down sorting method produced a production efficiency over the surface layer for both sorting parcels but also those in the mid to upper troposphere. Sorting will not only affect the reference state position of surface time. This suggests the restriction on surface CAPE release during further outward than the top-down results at the same integration anvil of parcels with positive APE production efficiency extended to upper troposphere. This is particularly obvious at 120 h as the anvil of parcels with positive APE production efficiency extended further outward than the top-down results at the same integration time. This suggests the restriction on surface CAPE release during sorting will not only affect the reference state position of surface parcels but also those in the mid to upper troposphere.

Figure 7(a) shows the time series of the area-averaged APE production efficiency over the surface layer for both sorting methods. In general, the top-down sorting method produced a much higher average surface APE production efficiency since more surface parcels were relocated upward and attained a smaller $T_{up}$ by releasing CAPE. After an initial spike up to 0.22 in the first 20 h, the average surface APE production efficiency reduced slowly to 0.15 at the end of the simulation. There were also signs of periodic fluctuation with a period of roughly 20–40 h.

Figure 7(b) shows the time series of the area-averaged top-down surface APE production efficiency over the inner and outer regions respectively. When averaging over the inner region only, the surface APE production efficiency increased steadily from 30 h onwards due to the build-up of CAPE by the continuous surface moisture flux. In contrast, the time series produced using data from the outer region only is characterised by a gradual decline from 30 h onward which is more consistent with the time series shown in Figure 7(a). The gradual reduction of the surface APE production efficiency in the outer region is related to the fluctuations in surface radial velocity described earlier in Figure 2. Figure 8 shows the radius–time plot of the top-down APE production efficiency in the lowest model level. The top-down surface APE production efficiency in the outer region dropped from above 0.35 to below 0.05 after each surface radial velocity fluctuation. This suggests the buoyancy of parcels in the outer region was reduced after the onset of each fluctuation which is consistent to the drop in $\theta_e$ shown earlier in Figure 2(b).

In contrast, the bottom-up sorting method produced a much lower average surface APE production efficiency due to the restriction on the release of surface CAPE during the sorting (Figure 7(a)). The average surface APE production efficiency remained fairly constant at 0.04 in the first 70 h and then increased gradually to 0.11 by the end of the simulation.

When averaging over the inner region only, the bottom-up surface APE production efficiency showed a rapid increase between 70 and 120 h and reached similar values to the top-down results by the end of the simulation. This increase was caused by the build-up of moisture in the boundary layer of the inner region which lowered the LCL of surface parcels such that they could be saturated by lifting them over a shorter distance. Although the bottom-up sorting method generally restricts the release of CAPE, when constructing the reference state the $P_{ref}$ slots nearer to the surface are usually assigned to the denser surface parcels from the outer region. As a consequence, surface parcels from the inner region can be ‘stacked’ above their original position when using the bottom-up sorting method. Combined with the lowering of LCL, a larger number of surface parcels were able to release CAPE towards the end of the simulation, resulting in a higher average surface APE production efficiency inside the inner region.

When averaging over the outer region only, the bottom-up surface APE production efficiency remained under 0.05 for the entire simulation and showed no clear trend of change. While the top-down results were affected by the outward propagation of surface radial velocity fluctuations, the bottom-up results were less sensitive. This is due to the restriction on CAPE release in the

Figure 7. Time series of the area-averaged top-down (blue) and bottom-up (green) APE production efficiency in the surface layer over (a) the whole domain, and (b) the inner region (radius ≤1000 km, solid lines) and outer region (radius >1000 km, dashed lines).

Figure 8. Radius–time plot of top-down production efficiency at the lowest model level.
bottom-up sorting method such that the reduction in buoyancy associated with the fluctuation will not have a significant impact on the bottom-up results especially in the outer region.

The results above showed that the surface APE production efficiency is controlled predominantly by the sorting method used to produce the reference state. The surface APE production efficiency also showed significant spatial variation between the inner and outer region, with the highest APE production efficiency always located within the inner region. The outward propagation of surface radial velocity fluctuations also affected the buoyancy of air parcels in the outer region. However, only the top-down results were sensitive to such a process. In the next section, the production of APE by surface fluxes is computed using the surface APE production efficiency.

### 3.3. Surface APE production

Using the APE production efficiency, the production of APE by surface fluxes is computed following Pauluis (2007). The production of APE by surface sensible and latent heat flux, $G_{\text{sen}}$ and $G_{\text{lat}}$ respectively, can be computed using

$$G_{\text{sen}} = T_1 - T_{1\text{ref}} \frac{Q_{\text{sen}}}{c_p \rho},$$

$$Q_{\text{sen}} = C_T (\theta_{\text{surf}} - \theta_{\Delta z/2}) \left( u^2 + v^2 \right)^{1/2} \Delta z/2;$$

$$G_{\text{lat}} = T_1 - T_{1\text{ref}} \frac{Q_{\text{lat}}}{L_v \rho},$$

$$Q_{\text{lat}} = C_E \left( r_{v,\text{surf}} - r_v \Delta z/2 \right) \left( u^2 + v^2 \right)^{1/2} \Delta z/2. \quad (8) \quad (9)$$

In Eq. (8), $Q_{\text{sen}}$ is surface sensible heat flux, $c_p$ is heat capacity and $C_T$ is the dimensionless flux coefficient for sensible heat. In Eq. (9), $Q_{\text{lat}}$ is surface latent heat flux, $L_v$ is latent heat of evaporation and $C_E$ is the dimensionless flux coefficient for moisture. The subscripts $\text{surf}$ and $\Delta z/2$ denote that the variable is evaluated at the surface and at a model half-level above the surface respectively. Both $G_{\text{sen}}$ and $G_{\text{lat}}$ have units of W m$^{-2}$.

The discussion here will focus on $G_{\text{lat}}$ since $G_{\text{sen}}$ is relatively insignificant. The radial distribution of surface latent heat flux within the vortex at both 60 and 120 h is consistent with the typical distribution of wind speed, with the largest values found inside the eyewall and a gradual reduction outwards from the inner region (Figure 9(a) and (b)). The radial profiles of $G_{\text{lat}}$ computed with the two sorting methods show a similar distribution, with the largest production found in the eyewall. Compared to the surface latent heat flux, the values of $G_{\text{lat}}$ are relatively small with roughly 30% of the surface latent heat flux converted into APE. The values of $G_{\text{lat}}$ are affected by the difference between the top-down and bottom-up APE production efficiency at the surface, which is most obvious at 45 h. When using the top-down APE production efficiency, positive values of APE production were found just outside the centre of rotation and extended radially outward to as far as 400 km radius. In contrast, little to no APE production was found beyond 50 km radius when using the bottom-up APE production efficiency. This is because the top-down APE production efficiency in the surface layer was generally larger than the bottom-up APE production efficiency especially in the outer region, as shown in section 3.2. The difference between the top-down and bottom-up APE production efficiencies in the surface layer became smaller in the inner region as the vortex intensified. At 120 h, the surface APE productions computed with both efficiencies were almost identical inside the inner region, although a slightly larger production was found between 800 and 1000 km radius with the top-down APE production efficiency.

Figure 10(a) shows the time series of surface latent heat flux and $G_{\text{lat}}$ computed using the top-down and bottom-up production efficiencies. The time series of both the surface latent heat flux and APE productions are computed for the inner region only, which better reflects the vortex’s intensification. The time series of surface latent heat flux showed little change in the first 40 h, followed by a sharp increase from just above 120 W m$^{-2}$ to more than 350 W m$^{-2}$ between 45 and 70 h as the surface wind speed strengthened during the rapid intensification. Upon reaching a steady state intensity, the surface latent heat flux remained consistently above 350 W m$^{-2}$ between 70 and 110 h before decreasing gradually to below 300 W m$^{-2}$ towards the end of the simulation. The reduction in surface latent heat flux near the end of the simulation was caused by the decreasing difference in $r_v$ between the boundary layer and the sea surface as well as a slower intensification of the surface wind speed. Both factors reduced the effectiveness of the surface latent heat flux.

Compared to the surface latent heat flux, the values of $G_{\text{lat}}$ in the time series were much smaller. While the radial profile (Figure 9) showed that as much as 30% of the surface latent heat flux can be converted into APE within the inner region, the time series of both APE productions generally show smaller percentages after including surface parcels with lower production efficiency at larger radii into the calculation when averaging out to 1000 km radius. Figure 10(b) shows a clearer illustration of both $G_{\text{lat}}$ profiles by reducing the range of the $y$-axis. When using the top-down APE production efficiency, the $G_{\text{lat}}$ profile was generally larger than that computed with the bottom-up APE production efficiency, which is consistent with the radial profile shown previously in Figure 9. The top-down $G_{\text{lat}}$ profile showed

![Figure 9. Radial profile of surface latent heat flux (SLHF dashed cyan) and $G_{\text{lat}}$ computed using the top-down (solid blue) and bottom-up (green) APE production efficiency at (a) 45 h and (b) 120 h. Note that the profiles extend to 1000 km radius only for clarity.](image-url)
a marked increase between 40 and 60 h as the vortex rapidly intensified. This was followed by a steady increase until 120 h and a slight decline near the end of the simulation.

Since the top-down surface APE production efficiency was affected by the fluctuations in surface radial velocity, the values of top-down $G_{\text{lat}}$ also showed similar variation. Figure 11 shows the radius–time plot of the values of $G_{\text{lat}}$ computed with the top-down APE production efficiency. Similar to the radial profile shown previously in Figure 9(a) and (b), the inner region within 1000 km radius was dominated by relatively large values of $G_{\text{lat}}$ while little to no $G_{\text{lat}}$ was found beyond 2500 km radius. The region between 1000 and 2500 km radius was characterised by alternating bands of higher (greater than 10 W m$^{-2}$) and lower (less than 10 W m$^{-2}$) $G_{\text{lat}}$ values. The transition from higher to lower values of $G_{\text{lat}}$ is quite rapid due to the rapid drop in the top-down APE production efficiency associated with fluctuations in surface radial velocity.

Compared to the top-down APE production time series, the bottom-up APE production time series mostly has smaller values, especially during the first 100 h of simulation. Also, the bottom-up APE production showed only a modest increase during the rapid intensification period between 40 and 60 h compared to the top-down APE production time series. The rise in bottom-up APE production began later at 70 h and showed a steady increase until 120 h, followed by a slight decline near the end of the simulation similar to the top-down APE production. The delayed rise of bottom-up APE production is consistent with the time series of the bottom-up APE production efficiency (Figure 7) which showed a much lower APE production efficiency in the first 70 h of the simulation followed by a gradual increase afterwards.

Finally, the production of APE by surface latent heat flux is compared to the conversion into KE ($C_{\text{APE→KE}}$). The conversion term $C_{\text{APE→KE}}$ is defined using the term representing buoyancy force in the vertical momentum equation:

$$C_{\text{APE→KE}} = \rho g \left( \frac{\partial \theta}{\partial z} + 0.61 (q_v - \overline{q_v}) - \overline{q_v} - C_{\text{F}} \frac{\partial \pi}{\partial z} \right). \quad (10)$$

In brief, the first and second terms on the RHS of Eq. (10) represent the contribution to buoyancy force from the deviation of $\theta$ and $q_v$ from the initial state $\overline{\theta}$ and $\overline{q_v}$ while the third term represents the work required to carry the liquid water loading. The last term is a ‘correction’ term needed to obtain the true buoyancy at a given integration time since the model domain will become warmer and more humid relative to the initial state as the vortex develops. This correction is made using the vertical gradient of the Exner pressure deviation term $\partial \pi_i/\partial z$ which is positive for an atmosphere that is warmer than the initial state thus lowering the buoyancy term and vice versa. A more detailed discussion on the role of buoyancy force and the correction term in the vertical KE budget is provided in the Appendix.

The time series of the conversion term, integrated over the volume of the inner region, is provided in Figure 10(b). In general, the conversion term increased gradually as the vortex intensified without any sharp increase during the rapid intensification period. This is because any increase in conversion during the rapid intensification period is likely to be very localised (e.g. inside the deep convections within the eyewall). Therefore, the conversion term time series in Figure 10(b), produced using the volume integral of the conversion term, is unlikely to pick up such localized increase during the rapid intensification period. Compared to the APE production time series, the bottom-up time series clearly shows a closer match to the conversion term, especially during the first 100 h of the simulation. This suggests that, when using the bottom-up sorting method, the APE budget was dominated by the balance between APE production and the conversion term while the $d\text{APE}/dt$ term was relatively small. However, the bottom-up $G_{\text{lat}}$ became greater than the conversion term in the last 30 h of the simulation. This was caused by the gradual increase in the bottom-up surface APE production efficiency especially in the inner region, as shown previously in Figure 7.

In contrast, the top-down APE production is always larger than the conversion term throughout the entire simulation, indicating
a build up of APE in the domain. Therefore, the \( \text{dAPE}/\text{d}t \) will be more significant in the APE budget due to the larger APE production resulting from the top-down sorting method.

The results above suggest the bottom-up sorting method is more capable in identifying the portion of APE generated by surface latent heat flux that can be converted into KE through the work of buoyancy forces. The top-down sorting method, in contrast, tends to generate much larger surface APE production as it assumes all surface CAPE in the vortex can be released, resulting in a larger production efficiency at the surface. However, a large portion of surface CAPE is dynamically inert, especially in regions dominated by subsidence further away from the centre of rotation. The large APE production in the top-down sorting method can only contribute to the storage term instead of converting into KE. Therefore, the conclusion here is that the bottom-up sorting method is a more suitable way to construct the reference state as it is able to minimize \( \text{dAPE}/\text{d}t \).

4. Conclusion and discussion

How to construct the reference state in moist APE theory has been a longstanding vexing issue. We investigated this here in connection with the energetics of TC intensification and MPI theory. Until now, the prevailing view had been that the reference state should be constructed as the one minimizing potential energy in an adiabatic re-arrangement of the fluid parcels, in accordance with Lorenz (1955)’s original recommendation. Although various authors such as Lorenz (1978), Randall and Wang (1992) and Tailleux and Grandpeix (2004) have discussed various ways to construct such a reference state, the implications for our understanding of moist energetics have been limited so far to rather abstract considerations about how to generalize the concept of CAPE in a one-dimensional atmosphere for which all the APE resides in its vertical component. Discussions of the implications of the choice of any particular reference state for our understanding of the energetics of concrete weather phenomena have been lacking.

In this study, two different sorting strategies were used to construct the reference state, referred to as the bottom-up and top-down sorting approaches, similarly to what was recently done for an ocean with a nonlinear equation of state by Saenz et al. (2015). In contrast to the oceanic case for which the two different approaches yield similar reference states, large differences exist between the bottom-up and top-down reference states in our simulation of TC intensification. This difference has important consequences for our understanding of the thermodynamic efficiency associated with the different diabatic processes at work in a TC and for trying to predict how much APE can contribute to the intensification.

As can be expected, the top-down reference state is the one that leads to the largest value of APE, since it is by construction the one that can incorporate most if not all the CAPE present in the domain, and therefore the one that the prevailing wisdom would recommend to use in moist APE theory. However, it is important to recognize that CAPE contributes to moist energetics only when parcels are able to reach their level of free convection, and hence when they have become absolutely unstable. The main problem with the top-down-based definition of APE is that it incorporates a large fraction of the total CAPE that is in fact dynamically inert and not actually available for conversion into KE. Many such parcels can be found in the regions of the TC dominated by subsidence, where CAPE is difficult if not impossible to release. In contrast, the bottom-up-based definition of APE only includes the CAPE of the absolutely unstable parcels, when such parcels have actually started to rise in updraughts. As a result, the top-down sorting method tends to overestimate the thermodynamic efficiency of the system, and hence to lead to an APE generation term that is much larger than the actual conversion of APE into KE. A much better agreement between APE generation and subsequent conversion into KE is found when using the bottom-up sorting method. Therefore, while the use of the top-down reference state might be preferable as a way to synthesize information about all available potential energy contained in the system, it nevertheless yields a less satisfactory description of the energetics of TC intensification for which it appears preferable to include as part of the APE definition only the part of the APE actually convertible into KE.

The present results have two important implications. First, they establish that the common approach of defining the reference state as one that minimizes the potential energy in an adiabatic re-arrangement may not be the most suitable approach, and that it is much better in the present example to define it such as to minimize \( \text{dAPE}/\text{d}t \) in order to maximize the correlation between APE generation and conversion of APE into KE; our results suggest that the bottom-up sorting approach allows one to achieve this objective. Second, they imply that we can provide a more rigorous footing for the MPI theory using APE theory; this will be investigated in future studies.

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Appendix

Buoyancy force and the vertical kinetic energy budget

The key result presented in this article is that given a suitable reference state, the production of APE by surface latent heat flux is mostly balanced by the conversion into KE through the work of buoyancy force. Such conversion takes place via the vertical momentum equation, which in the 2D model is given by

\[
\frac{Dw}{Dt} = g \left( \frac{\theta - \bar{\theta}}{\bar{\theta}} + 0.61 (\theta - \bar{\theta}) - \bar{q}_l \right) - c_p \bar{\theta} \frac{\partial \bar{\theta}}{\partial z} + D_w, 
\]

where \( w \) is vertical velocity, \( g \) is gravity, \( \theta \) and \( \bar{\theta} \) are the initial potential temperature and vapour mixing ratio profiles used to initialize the model domain, \( q_l \) is liquid water mixing ratio, \( c_p \) is heat capacity, \( \bar{\theta} \) is the initial virtual potential profile, \( \pi \) is the Exner pressure deviation from the initial pressure profile and \( D_w \) is diffusion of KE. Using the vertical momentum equation, the vertical KE budget can be defined as

\[
\frac{DKE_{w}}{Dt} = wmg \left( \frac{\theta - \bar{\theta}}{\bar{\theta}} + 0.61 (\theta - \bar{\theta}) - \bar{q}_l \right) - mw \pi \frac{\partial \pi}{\partial z} + w D_{w} ,
\]

where \( KE_{w} = 0.5 m w^2 \) is the vertical KE and \( m \) is mass. Equation (A2) shows that the vertical KE budget is controlled by three terms on the right-hand side that represent the work of buoyancy force, the work of vertical PGF and diffusion of vertical KE respectively. Note that the work of buoyancy force in Eq. (A2) is defined using the deviation of \( \theta \) and \( \bar{\theta} \), from the initial profiles \( \bar{\theta} \) and \( \pi \). Both deviation terms are generally positive due to the continuous heating and moistening of the model domain as the vortex intensified. Therefore, the work of buoyancy force is a source of \( KE_{w} \) for an ascending air parcel with positive \( w \). Meanwhile, \( \partial \pi/\partial z \) is usually positive due to the less rapid drop of air pressure in the mid-atmosphere compared to the surface. Therefore, the work of vertical PGF and the diffusion of vertical KE are net sinks of \( KE_{w} \) for an ascending air parcel.

Since the work of buoyancy force is clearly a source of \( KE_{w} \), it is therefore a suitable representation of the portion of APE that

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is converted into vertical KE (i.e. the conversion term $C_{APE\rightarrow KE}$). However, a key problem here is that the work of buoyancy force in Eq. (A2) is defined with respect to the $\bar{\theta}$ and $\bar{q}$ profiles which are time independent while the reference state and associated APE production efficiency evolve with time. In order to link the conversion term to the time-dependent reference state and the APE production term, the work of buoyancy force in Eq. (A2) is redefined with respect to a time-dependent horizontally averaged Exner pressure field. The Exner pressure deviation ($\pi$) is divided into a time-dependent horizontal average across the vortex ($\pi_r$) and a local deviation ($\pi'$) such that $\pi = \pi_r + \pi'$. The vertical KE budget can then be written as

\[ \frac{DKE_w}{Dt} = \text{wmg} \left( \frac{\theta - \bar{\theta}}{\bar{\theta}} + 0.61(q_r - \bar{q}) - q_l - \frac{\epsilon p_s}{g} \frac{\partial \pi_r}{\partial z} \right) - mwc_p \frac{\partial \pi'}{\partial z} + \text{wmd}_w. \tag{A3} \]

The $\frac{\partial \pi_r}{\partial z}$ term in the first term on the RHS of Eq. (A3) acts as a correction that removes the extra buoyancy with respect to the time-independent initial profiles. By incorporating it into the term representing the work of buoyancy force (i.e. first term on the RHS of Eq. (A3)), it can then be used for a fair comparison with the time-dependent APE production term as in Eq. (10).

References


