Matter Density versus Distance for the Neutrino Beam from Fermilab to Lead, South Dakota, and Comparison of Oscillations with a Variable and a Constant Density

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This paper is divided into two parts. In the first part, the material densities passed through for neutrinos going from FNAL to Sanford Laboratory are calculated using two recent density tables, Crustal [5] and Shen-Ritzwoller [4]. The method of calculation here can be used as a template for finding the densities along other long paths, as well as the values from an older table PEMC [7]. For the programs and tables, see the website: http://igppweb.ucsd.edu/~gabi/crust1.html and http://ds.iris.edu/ds/products/emc-pem/

In the second part of this paper, oscillations calculated using two recent density tables, Crustal [5] and Shen-Ritzwoller [4], as well as the values from an older table PEMC [7], are compared with oscillation results using the mean density from the Shen-Ritzwoller tables and one other fixed density. For the tests made here, the mean density results are quite similar to those found using the variable density vs distance.

I. INTRODUCTION

The Long-Baseline Neutrino Facility (LBNF) [1] and the Deep Underground Neutrino Experiment (DUNE) [2], now under preliminary construction will send a beam of neutrinos from the Fermi National Accelerator Laboratory (FNAL) near Chicago to the Sanford Laboratory located in a former gold mine in Lead, South Dakota. The neutrino beam will travel through varying densities of material along its path. Along its way the neutrinos will oscillate between the three known kinds of neutrinos. This oscillation is affected by the presence of the material or, more precisely, by the density of electrons along its path [3]. Although it is possible to calculate the oscillations expected on a variable density path, most of the preliminary calculations have assumed a constant average density. An early LBNF report [4] stated that to include the effects of variable density, a 5% density systematic error.

In the first part of this paper, the variable density travelled by the neutrinos along their path is calculated using two recent density tables, Crustal [5] and Shen-Ritzwoller [4], as well as the values from an older table PEMC [7]. The method of calculation here can be used as a template for finding the densities along other long neutrino beams.

II. FINDING DENSITIES ALONG THE NEUTRINO PATH

A. Dividing Up The Path

The earth is approximately an ellipsoid [8]. The radius in the polar direction is 6356 km and in the equatorial direction is 6378 km. Both of these numbers are accurate to better than 0.1 km.

Twenty five points were selected taking equal intervals of latitude (lat) and longitude. For two points at the same latitude, the distance between the two longitude points is not constant, but varies as cos(lat) going from zero at the poles to a maximum at the equator. For the DUNE beam path, the adjacent points have slightly different latitudes. However, the latitude differences between adjacent points are quite small and taking a mean value between adjacent points introduces a negligible error.

Let the distance from the center of the earth to sea-level at a given latitude-longitude value be \( RL_i \), the local radius at point \( i \). For \( i > 1 \) let \( \Delta \theta_i \) be the angle between \( RL_i \) and \( RL_{i-1} \), and \( \theta_i \) be the total angle between the initial local radius (\( RL_1 \) and \( RL_i \)).

\[
(x/6378)^2 + (y/6356)^2 = 1. \tag{1}
\]

Then \( x_i = RL_i \cos(\text{lat}) \); \( y = RL_i \sin(\text{lat}) \).

\[
1/RL_i = \sqrt{(\cos(\text{lat})/6378)^2 + (\sin(\text{lat})/6356)^2}. \tag{2}
\]

If we have a flat earth then then we would go from the initial height to final height linearly with distance (dist(\( i \))) along the neutrino beam. Let \( \text{ftosl} \) be the distance along the neutrino beam from FNAL to Sanford Laboratory.

\[
\text{flat height}(i) = (\text{endsseaheight} * \text{dist}(i) + \text{startseaheight} * (\text{ftosl} – \text{dist}(i))/\text{ftosl}. \tag{3}
\]

In the second part of this paper, oscillations calculated using the variable density path are compared with two fixed density calculations.
The start height of the beam at FNAL is 228.4 m above sea level and the end height of the midpoint of the detector at Sanford Laboratory is 159 m.

For the curved earth part starting and ending at sea-level with a total arc of $\theta_{\text{total}}$, the angle of the arc is taken from $-\theta_{\text{total}}/2$ to $+\theta_{\text{total}}/2$. For 25 points, the midpoint in the neutrino beam path would be given by point 13 if the latitude of FNAL and Sanford Laboratory were the same. In fact, they are at different latitudes and that introduces a non-symmetric change in the path segment lengths along the beam path, which changes the center point slightly. Empirically it is found to be located 2% of the way between point 13 and point 14, “point 13.02”.

See Fig. 1. Let $L$ be the straight line connecting the sea level points at initial and final destinations, $R$ be the local radius at the center of the beam path, $s$ be the perpendicular distance from the midpoint of $L$ to the circle (the sagitta), and $t$ be the distance along the local radius from a point on $L$ at a distance $d$ from the start to the local circle.

\[
R^2 = (R - s)^2 + (L/2)^2; \\
(R - s)^2 = R^2 - (L/2)^2. \tag{4}
\]

$t$ is not quite perpendicular to the straight line $L$, but the error is small. The fractional error in $t$ is zero at the center of the arc and increases, approximately quadratically, approaching a value of 0.5% of the perpendicular distance by the end of the arc, where $t$ is very small.

\[
(R - s)^2 + (d - L/2)^2 = (R - t)^2. \tag{5}
\]

Substitute Equation 4 into Equation 5.

\[
R^2 - (L/2)^2 + (d - L/2)^2 = (R - t)^2; \\
(L/2)^2 - (d - L/2)^2 = 2Rt - t^2. \tag{6}
\]

Ignore the $t^2$ term.

\[
t = [(L/2)^2 - (d - L/2)^2]/(2R). \tag{7}
\]

For the calculation of $t$, the variation of the local radius over the path segment from $i$ to $i + 1$ is produces a negligible effect. The distance above sea level at distance $d$ is then given by the sum of the flat height and the curved height ($t$). There is an additional effect called the geoid height [9], but it is very small, about 0.01 m for the FNAL point and −13.7 m for the Sanford Laboratory point.

Let $\theta_{\text{midpoint}}$ be the angle between the local radius for point 1, and the midpoint radius. For point $i$, the angle that $t$ makes with the midpoint radius is $\theta_i - \theta_{\text{midpoint}} = \alpha$. This angle is also the angle that the tangent to the local radius circle makes with the line $L$. For this short segment the length of the arc and the length of the chord are essentially equal.

For $i > 1$, the straight line distance from FNAL to Sanford Laboratory is incremented by

\[
dist(i) = dist(i-1) + \cos(\alpha) \times RL_i \times \Delta \theta_i. \tag{8}
\]

The distance from FNAL to Sanford Laboratory seen by the neutrino beam (fltosl) is calculated to be fltosl = 1284.9 km.

The density maps depend on the depth of the beam below ground at the various points. At Sanford Laboratory there are a number of hills and the beam ends up above sea level even though the center of the detector is close to 1470 m beneath the surface. The elevation at a given latitude and longitude can be obtained from a convenient web site [10] and the difference between the elevation and the sea level height of the beam is then the depth. See Fig. 2. In general the elevation varies smoothly except very near to Sanford Laboratory. If the elevation had fluctuated considerably over a fair fraction of the path it would have added uncertainty to the density map.

B. Results And Their Uncertainties

Crustal is a recent (2013) attempt to find the density of the earth as a function of latitude and longitude. CRUST1.0 is an 8 layer model. Although it is not needed here, a ninth layer gives the density below the Moho. Crustal averages crust structure over 1 x 1 degree cells (about 110 x 110 km). The map is based on the ETOP01 global relief model produced by the National Centers for Environmental information, a part of the National Oceanic and Atmospheric Administration[11].

The model is defined from +89.5 to −89.5 deg. lati-
tude and −179.5 to +179.5 deg. longitude. Density is in gm/cm$^3$. Our longitude (W) corresponds to negative values here. Crustal supplies a program (getCN1point) which for a given latitude and longitude at the midpoint of a cell, gives the density of each layer and the bottom of the layer. For all maps in this paper, the depth, not the sea-level height is used in the maps.

The Shen-Ritzwoller model is a new (2016) density map only of the United states in $1/4 \times 1/4$ degree cells of latitude and longitude. The density map is divided into many more layers, than the Crustal map. There are more than 50 layers.

There is also an older map, PEMC included for historical reasons. A comparison of the density vs distance results of each map is shown in Fig. 3 and the numerical results are given in Tables I and II.

Although the actual situation is more complicated, we will look at uncertainties in the total amount of matter passed through by the neutrinos ($\int \rho dx$) to get an indication of uncertainties. There are two kinds of uncertainties to be considered, statistical and systematic. Statistical uncertainties are due to random differences. Sometimes the depths are near a boundary between two densities. The boundaries are probably not completely flat and there is some transition region. In the crustal map there are six points within about 1.5 km of a depth boundary with an average change in density of about 4%. If we view this as a random walk then the standard deviation in the total amount of matter passed through is 0.43%. Even if all twenty-five path segments had a 4% uncertainty, the standard deviation in the total amount of matter passed through would be 0.8%. The statistical uncertainties are quite small.

There are many more layers given for the Shen-Ritzwoller map and the differences from layer to layer...
are of the order of 1% (except for the last point, which has 15% differences). The statistical uncertainties are again small.

The systematic uncertainties are those due to a systematic error in the density of the layers. One approach is to compare the mean density for the three maps. The mean density for PEMC is 2.845 gm/cm$^3$ for Crustal it is 2.817 gm/cm$^3$ and for Shen-Rittzower it is 2.848 gm/cm$^3$. The PEMC map and the Shen-Rittzower map have essentially identical means while the Crustal mean is approximately 1% lower.

Some early DUNE calculations used a mean density of 2.957 gm/cm$^3$ and a distance of 1300 km [12]. This density is 4% higher than the Shen-Rittzower mean density and 5% higher than the crustal mean density. In addition, the distance is 1% longer than the distance calculated here (1284.9 km), so the total amount of material through which the beam passes is 5% or 6% higher than the numbers here.

For the Shen-Rittzower map there is another way to estimate errors. They are still calculating detailed systematic errors, but they suggest that a reasonable estimate of the error in density is to use the standard deviation in shear velocity ($v_s$) given in their Fig. 15 together with the empirical relation between $v_s$ and $\rho$ obtained by T.M. Brocher [13],
\[
\rho = 1.227 + 1.53v_s - 0.837v_s^2 + 0.207v_s^3 - 0.01066v_s^4.
\]

In their Fig. 15, the standard deviation in the magnitude of $v_s$ is of the order of 0.03 to 0.05 km/sec over the region of the DUNE beam. The fractional errors in density obtained are fairly constant over the beam path. For 0.03, 0.05, and 0.07 km/sec errors in $v_s$, one obtains mean fractional errors in density of 0.5%, 0.8%, and 1.2%.

### C. Electron Density Distribution In The Earth

For a single kind of atom with atomic number $Z$ and given atomic weight, the number of atoms in one gm-atomic weight is Avagadro’s number ($N_{Av}$). Let $\rho$ = the density of the material in gm/cm$^3$. The number of electrons in one cubic centimeter ($N_e$) is then
\[
N_e = Z \times N_{Av} \times \rho/\text{atomic wgt.}
\]

For a mix of materials the quantity needed is the mean value of $Z$/atomic wgt. Tables of the abundance in parts per million (ppm) of the various elements in the crust are given in reference [14]. In fact this reference lists three tables of abundances [15–17]. The tables are in reasonable agreement for the main components, but some of the minor elements differ by 20% or more. Table III gives the abundances for the most abundant 9 elements, in ppm. The further elements are present only at the level of < 0.3%. (Fe is the most abundant element at lower depths, but not at the depths appropriate to this beam.)

| Element | Z | Table 15 | Table 16 | Table 17 | Mean |
|---------|---|----------|----------|----------|------|
| O       | 8 | 460000.  | 467100.  | 461000.  | 462700.|
| Si      | 14 | 270000.  | 276900.  | 282000.  | 276300.|
| Al      | 13 | 82000.   | 80700.   | 82300.   | 81667. |
| Fe      | 26 | 63000.   | 50500.   | 56300.   | 56600. |
| Ca      | 20 | 50000.   | 36500.   | 41500.   | 42667. |
| Na      | 11 | 23000.   | 27500.   | 23600.   | 24700. |
| K       | 19 | 15000.   | 25800.   | 20900.   | 20567. |
| Mg      | 12 | 29000.   | 20800.   | 23300.   | 24367. |
| Ti      | 22 | 6600.    | 6200.    | 5600.    | 6133.  |

TABLE III. Abundances in ppm of the major elements in the Earth’s crust.

In addition the abundance of stable isotopes and atomic weights of these nine elements are needed [13].
TABLE IV. Isotopic numbers (A) and isotopic weights of stable isotopes of the major elements in the Earth’s crust.

| El. | A wgt | A wgt | A wgt | A wgt | A wgt | A wgt |
|-----|-------|-------|-------|-------|-------|-------|
| O   | 16    | 17    | 16    | 17    | 16    | 17    |
| Si  | 28    | 29    | 28    | 29    | 28    | 29    |
| Fe  | 54    | 56    | 55    | 56    | 54    | 56    |
| Ca  | 40    | 42    | 41    | 43    | 40    | 42    |
| Na  | 23    | 25    | 24    | 26    | 23    | 25    |
| K   | 39    | 41    | 40    | 42    | 39    | 41    |
| Mg  | 24    | 26    | 25    | 27    | 24    | 26    |
| Ti  | 46    | 48    | 47    | 49    | 46    | 48    |

TABLE V. Percentage isotopic abundances of stable isotopes of the major elements in the Earth’s crust.

| El. | A abund | A abund | A abund | A abund | A abund |
|-----|---------|---------|---------|---------|---------|
| O   | 99.757  | 0.038   | 0.205   |         |         |
| Si  | 92.230  | 4.683   | 0.087   |         |         |
| Fe  | 100.0   |         |         |         |         |
| Ca  | 96.941  | 0.647   | 2.086   |         |         |
| Na  | 100.0   |         |         |         |         |
| K   | 93.258  | 6.730   |         |         |         |
| Mg  | 78.99   | 10.0    | 11.01   |         |         |
| Ti  | 8.25    | 7.44    | 73.72   | 5.41    | 5.18    |

The atomic weights are given in Table IV and the percentage fractional isotopic abundances in Table V. Table VI gives $Z$/atomic weight and $Z$/A averaged over the elements for each of the three abundance tables and for the mean, as well as the standard deviation from the three tables.

For the mean abundance, the number of electrons per cubic centimeter for $\rho = 1$ is $2.9805 \times 10^{23}$. The fact that $Z/A$ is so near to 1/2 is not surprising. The most abundant elements, oxygen (O) and silicon (Si), comprising about 75% of the total have isotopic abundances overwhelmingly favoring 1/2.

| $Z$/wgt | Mean | $\sigma$ |
|---------|------|---------|
| 0.4948  | 0.4945 | 0.4945 |
| 0.4949  | 0.4949 | 0.4949 |

TABLE VI. Average $Z$/atomic weight, and $Z$/A, using the three different abundance tables. The fourth column is the result for the mean abundance from the three tables and the fifth column is the standard deviation of the three values.

III. NEUTRINO OSCILLATION PROBABILITIES AT SANFORD LABORATORY

For the present analysis, the density results using the new Shen-Ritzwoller map are used with one small modification. It was more convenient to have the neutrino beam distances between points constant for the density vs distance map. Here the average distance between points was used. The maximum distance change was about 6 km. That occurred at the center of the path, where the density changes from point to point are small.

Neutrino oscillations are calculated for a variable density path using the computer program of J. Kopp [19]. Results are presented for this variable density map, for a constant density of 2.848 gm/cm$^3$, which is the mean density for this variable density map, and for the density of 2.957 gm/cm$^3$. The distance between FNAL and the Sanford Laboratory was calculated in Section II to be 1284.9 km. The DUNE calculations which used 2.957 gm/cm$^3$ used 1300 km as a distance. For the present comparison a distance of 1284.9 km was used for this density as well.

A. Plots of Oscillation Probabilities for the Variable Density Option

Figures 4–7 show plots of oscillation probabilities at Sanford Laboratory for $\nu$ and $\bar{\nu}$ oscillations separately, for both the CP violation parameter $\delta_{CP} = 0$ and $\delta_{CP} = 3\pi/2$. The differences between the three density options, for $\nu$ and $\bar{\nu}$, for $\delta_{CP} = 0$ and $\delta_{CP} = 3\pi/2$ have been calculated. As an example, the differences between the variable density option and the fixed 2.848 gm/cm$^3$ density option for $\nu$ and $\bar{\nu}$ with $\delta_{CP} = 0$ are shown in Figs. 8 and 9. Note the difference of probability scales between Figs. 4 to 7 and Figs. 8-9.

For Nu with no CP violation, variable density

FIG. 4. Pr($\nu$) oscillations with $\delta_{CP} = 0$ using the variable density path.

![Graph showing oscillation probabilities](image-url)
For Nu with no CP violation, variable minus fixed density of 2.848 gm/cm$^3$

For Nu with no CP violation, variable density

For Nu with CP violation, variable density

For Nubar with no CP violation, variable density

For Nubar with CP violation, variable density

dependent on the kind and range of the average. For the average will likely reduce the differences and be very practice energy bands will have to be selected. However, included only to give a sequence of energies encompassing

$\delta \ll 1$.

In Table VII, for the variable density assumption, three density assumptions, and $\Delta_1$ serves to emphasize the matter interactions.

The second quantity examined was:

$$\Delta_2(E) = (\text{Pr}(\nu_e) - \text{Pr}(\bar{\nu}_e)) \text{ for } \delta_{CP} = 0$$

$$- (\text{Pr}(\nu_e) - \text{Pr}(\bar{\nu}_e)) \text{ for } \delta_{CP} = 3\pi/2.$$  \hfill (12)

This is an important quantity to use to look at CP violation.

$\Delta_1(E)$ and $\Delta_2(E)$ were examined for each of the three density assumptions, and $\Delta_1(E)$ was examined both for $\delta_{CP} = 0$ and for $\delta_{CP} = 3\pi/2$.

In Table VII, for the variable density assumption, three energies corresponding to probability maxima for $\Delta_1$ and $\Delta_2$ are shown along with their maximum values.

For the following tables, “v” refers to the variable density assumption, “s” refers to a fixed density of $\rho = 2.848$
gm/cm³, and “d” refers to a fixed density of ρ = 2.957

gm/cm³. “v − s” means variable density minus fixed
density 2.848 gm/cm³, “v − d” means variable density
minus fixed density 2.957 gm/cm³, and “d − s” means
fixed density 2.957 gm/cm³ minus fixed density 2.848

gm/cm³. For comparisons involving the variable density,
the energies correspond to the three energy values in Ta-
ble VII. For the comparisons of “d − s” the values for
the peak energies for “d” nearest to those in Table VII were
chosen.

Table VIII examines the differences between the
Δ₁(Epeak₁) values for the different density assumptions,
where Epeak₁ is the energy of the maximum Δ₁ for the
first density assumption. δ(Δ₁(Epeak₁)) is the difference
of Δ₁ found in the two density assumptions. The per-
centages of the ratio δ(Δ₁(Epeak₁))/Δ₁(Epeak₁) are shown
for each of the three energies.

In Tables IX and X, Emax₁ is the nearest energy to
Epeak₁ for which |δ(Δ₁(Emax₁))| is at a local maximum.
The percentage differences of Emax₁ from Epeak₁ and of
the ratio δ(Δ₁(Emax₁))/Δ₁(Emax₁) are shown. Table IX
shows these quantities if δCP = 0 and Table X shows
these quantities if δCP = 3π/2. The Δ₁ differences are
sometimes appreciable, although the values of Δ₁ often
are small.

Table XI examines the differences between the
Δ₂(Epeak₂) values for the different density assumptions.
Epeak₂ is the energy of the maximum Δ₂ for the first
density assumption. δ(Δ₂(Epeak₂)) is the difference
of Δ₂ found in the two density assumptions. The per-
centages of the ratio δ(Δ₂(Epeak₂))/Δ₂(Epeak₂) are shown
for each of the three energies.

In Table XII, Emax₂ is the nearest energy to Epeak₂
for which |δ(Δ₂(Emax₂))| is at a local maximum. The
percentage differences of Emax₂ from Epeak₂ and of the
ratio δ(Δ₂(Emax₂))/Δ₂(Emax₂) are shown.

For Δ₂, the difference between using the variable den-
sity and the mean of the variable density, 2.848 gm/cm³
is small, of the order of 0.2%, except for the one anom-
alous value. That value occurs because the largest value

TABLE VII. Results for the variable density option for Δ₁ =
Pr(ν) − Pr(ν̄) and Δ₂ = Δ₁(δCP = 0) − Δ₁(δCP = 3π/2). The
columns labelled Δ are Δ₁ or Δ₂ as designated in column 1.
E is the energy of the chosen maximum Δ in GeV.

| Δ | δCP | E | Δ | E | Δ | E | Δ |
|---|-----|---|---|---|---|---|---|
| Δ₁ | 0 | 0.996 | 0.14 | 0.42 | -0.0521 | 1.12 | -0.028 |
| Δ₁ | 1.5π | 0.096 | 0.41 | 0.37 | 0.16 | 0.811 | 0.086 |
| Δ₂ | 0.096 | -0.27 | 0.37 | -0.15 | 0.827 | -0.069 |

TABLE VIII. Δ₁(Epeak₁) = Pr(ν) − Pr(ν̄). Epeak₁ is the
energy of the maximum Δ₁ for the first density assump-
tion. δ(Δ₁(Epeak₁)) is the difference of Δ₁ found in the
two density assumptions. The percentages of the ratio
δ(Δ₁(Epeak₁))/Δ₁(Epeak₁) are shown for each of the three
energies.

| Δ | δCP | δ(Δ₁) | δ(Δ₁) | δ(Δ₁) |
|---|-----|-------|-------|-------|
| v − s | 0 | 0.92 | 0.54 | 2.47 |
| v − d | 0 | -2.6 | -2.8 | 0.68 |
| d − s | 0 | 3.4 | -3.15 | |
| v − s | 3π/2 | 0.27 | 0.18 | 0.57 |
| v − d | 3π/2 | 0.21 | 0.007 | 1.0 |
| d − s | 3π/2 | 0.48 | -0.18 | 0.47 |

TABLE IX. Epeak₁ is the energy of the maximum Δ₁ for the
first density assumption and Emax₁ is the nearest energy to
Epeak₁ for which |δ(Δ₁(Emax₁))| is at a local maximum. The
percentage differences of Emax₁ from Epeak₁ and of the ra-
tio δ(Δ₁(Emax₁))/Δ₁(Emax₁) are shown for each of the three
energies. δCP = 0 is assumed for this table.

| Δ | δE | δ(Δ₁) | δ(Δ₁) | δ(Δ₁) |
|---|-----|-------|-------|-------|
| v − s | 0 | 0.92 | -4.9 | -0.91 | -7.8 | 3.5 |
| v − d | 0 | -2.6 | 0.47 | -2.8 | 6.7 | 0.98 |
| d − s | 0 | 0.92 | 3.4 | -0.97 | 3.16 |
TABLE X. $E_{\text{peak}1}$ is the energy of the maximum $\Delta_1$ for the first density assumption and $E_{\text{max}1}$ is the nearest energy to $E_{\text{peak}1}$ for which $|\delta(\Delta_1(E_{\text{max}1}))|$ is at a local maximum. The percentage differences of $E_{\text{max}1}$ from $E_{\text{peak}1}$ and of the ratio $\delta(\Delta_1(E_{\text{max}1}))/\Delta_1(E_{\text{max}1})$ are shown for each of the three energies. $\delta_{CP} = 3\pi/2$ is assumed for this table.

| $\Delta$ | $\text{var}$ | $dE$ | $\delta(\Delta_1)$ | $dE$ | $\delta(\Delta_2)$ | $dE$ | $\delta(\Delta_3)$ |
|----------|-------------|------|------------------|------|------------------|------|------------------|
| $v - s$  | -0.31       | 4.0  | 3.3              | 0.25 | 15.2             | 1.5  |
| $v - d$  | -3.1        | 10.1 | 8.4              | 1.2  | -9.0             | 1.3  |
| $d - s$  | -3.1        | 16.6 | 7.4              | 1.3  | -9.0             | 1.1  |

TABLE XI. $\Delta_2(E_{\text{peak}2}) = \Delta_1(\delta_{CP} = 0) - \Delta_1(\delta_{CP} = 3\pi/2)$. $E_{\text{peak}2}$ is the energy of the maximum $\Delta_2$ for the first density assumption. $\delta(\Delta_2(E_{\text{peak}2}))$ is the difference of $\Delta_2$ found in the two density assumptions. The percentages of the ratio $\delta(\Delta_2(E_{\text{peak}2}))/\Delta_2(E_{\text{peak}2})$ are shown for each of the three energies.

| $\Delta$ | $\text{var}$ | $dE$ | $\delta(\Delta_1)$ | $dE$ | $\delta(\Delta_2)$ | $dE$ | $\delta(\Delta_3)$ |
|----------|-------------|------|------------------|------|------------------|------|------------------|
| $v - s$  | -0.05       | -0.62| -0.31            |
| $v - d$  | 0.95        | 0.40 | 0.20             |
| $d - s$  | -1.0        | -0.53| -0.37            |

TABLE XII. $E_{\text{peak}2}$ is the energy of the maximum $\Delta_2$ for the first density assumption and $E_{\text{max}2}$ is the nearest energy to $E_{\text{peak}2}$ for which $|\delta(\Delta_2(E_{\text{max}2}))|$ is at a local maximum. The percentage differences of $E_{\text{max}2}$ from $E_{\text{peak}2}$ and of the ratio $\delta(\Delta_2(E_{\text{max}2}))/\Delta_2(E_{\text{max}2})$ are shown for each of the three energies.

| $\Delta$ | $\text{var}$ | $dE$ | $\delta(\Delta_1)$ | $dE$ | $\delta(\Delta_2)$ | $dE$ | $\delta(\Delta_3)$ |
|----------|-------------|------|------------------|------|------------------|------|------------------|
| $v - s$  | 4.0         | -0.794| -0.62            | -0.13| -4.0             | -0.20|
| $v - d$  | 6.3         | 1.1  | 1.6              | 0.48 | 9.7              | 0.53 |
| $d - s$  | 6.3         | 1.2  | 1.4              | -0.59| 6.7              | -0.56|

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