Standardizing anthropometric measures in children and adolescents with new functions for  
\texttt{egen}

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\textbf{Abstract.} A new function for  \texttt{egen}  has been developed to allow transformation of child anthropometric data to z-scores using the LMS method and the reference data available from the 1990 British Growth Reference and the 2000 US CDC Growth Reference. An additional function allows for children to be categorized according to body mass index (weight/height$^2$) using international cutoff points recommended by the Childhood Obesity Working Group of the International Obesity Taskforce.

\textbf{Keywords:} dm0004, z-scores, LMS,  \texttt{egen}

\section{Introduction}

Comparing anthropometric data from children of different ages is complicated by the fact that children are still growing. (We do not expect the height of a 5-year-old to be the same as the height of a 10-year-old!) Clinicians and researchers are often interested in the question, “Is this child taller, shorter, or about average compared to other children of the same age?” Two sets of population-based reference data are now widely used to address this question: the 1990 British Growth Reference and the 2000 Centers for Disease Control and Prevention (CDC) Growth Reference in the United States. The British Growth Reference was developed using data from seventeen cross-sectional surveys conducted in the United Kingdom between 1978 and 1993 (Cole, Freeman, and Preece 1998). The CDC Growth Reference was developed using data from five national surveys conducted in the United States between 1963 and 1994 and from supplemental birth data collected from birth certificates and medical records. The CDC Growth Reference has recently been updated (Kuczmarski et al. 2000; \url{www.cdc.gov/growthcharts}). Both references tabulate values obtained by the LMS method, developed by Cole (1990) and Cole and Green (1992), that can be used to transform crude data to standard deviation z-scores. These are standardized to the reference population for the child’s age and
sex (or for height/length and sex). The LMS transformation removes skewness and adjusts for physiological changes in anthropometric measures that occur with age. Stata provides a convenient environment in which to apply the age-specific (or height/length-specific) LMS values and generate z-scores for each child in a dataset, using the `egen` command.

Until recently, there was no standard definition of overweight and obesity for children. The Childhood Obesity Working Group of the International Obesity Taskforce recently recommended the use of body mass index (BMI) cutoff points to categorize children as normal weight, overweight, or obese based on age, gender, and BMI (Cole et al. 2000). These cutoff points were developed using large, nationally representative cross-sectional datasets from six countries: Brazil, Great Britain, Hong Kong, the Netherlands, Singapore, and the United States. They correspond to equivalent adult BMI cutoff points endorsed by the World Health Organization of BMI<25 kg/m² for normal weight, BMI 25–29.99 kg/m² for overweight, and BMI≥30 kg/m² for obesity. An additional function of `egen` categorizes children as normal weight, overweight, or obese according to these international cutoff points.

### 2 Syntax

```stata
egen [type] newvar = zanthro(varname, chart, version) [if exp] [in range],
   xvar(varname) gender(varname) gencode(male=code, female=code)
   [ageunit(unit) nocutoff ]
```

```stata
egen [type] newvar = zbmicat(varname) [if exp] [in range], xvar(varname)
   gender(varname) gencode(male=code, female=code) [ageunit(unit) ]
```

by ...: may not be used with either of these functions.

### 3 Functions

`zanthro(varname, chart, version)` calculates z-scores for anthropometric measures in children and adolescents according to US or UK reference growth charts. The three arguments are

- `varname`: the variable name of the measure in your dataset for which z-scores are to be calculated (e.g., `height`, `weight`, or `bmi`).
- `chart`: a valid chart code (see the tables in section 5 for a list of valid codes).
- `version`: either US or UK. Some charts are available for only one of these versions.

`zbmicat(varname)` categorizes children from 2 to 18 years as normal weight, overweight or obese, using BMI cutoffs recommended by the Childhood Obesity Working Group of the International Obesity Taskforce. BMI is in kg/m². This function generates a
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variable with values 1, 2, and 3 labeled, respectively, as “Normal wt”, “Overweight”, and “Obese”.

4 Options

\texttt{xvar(varname)} specifies the variable used (along with gender) as the basis for standardizing the measure of interest. This variable is usually age but can also be height or length when the measurement is weight; i.e., weight-for-age, weight-for-height, and weight-for-length are all available growth charts.

\texttt{gender(varname)} specifies the gender variable. It can be string or numeric. The codes for male and female must be specified by the \texttt{gencode()} option.

\texttt{gencode(male=code, female=code)} specifies the codes for male and female. The gender can be specified in either order, and the comma is optional. No quotes are required around the codes, even if the gender variable is a string.

\texttt{ageunit(unit)} gives the unit for the \texttt{age} variable and is only valid for measurement-for-age charts; i.e., omit this option when the chart code is \texttt{wh} or \texttt{wl} (see section 5). The \texttt{unit} can be \texttt{day, month, or year}. This option can be omitted if the \texttt{unit} is \texttt{year}, as this is the default.

\texttt{nocutoff} is optional. By default, any \texttt{z} scores with absolute value greater than or equal to 5 (i.e., values that are 5 standard deviations or more away from the mean) are set to missing. This option forces calculation of all \texttt{z} scores, allowing for extreme values in your dataset.

The decision to have a default cutoff at 5 standard deviations from the mean was made as a way of attempting to capture extreme data entry errors. Apart from this and setting to missing any \texttt{z} scores where the measurement is a nonpositive number, these functions will not automatically detect data errors. Please check your data, as always!

5 Growth charts

For both CDC and British charts, each growth reference is summarized by three numbers, called L, M, and S, which represent the skewness, median, and coefficient of variation of the measurement as it changes with \texttt{xvar()}. Where \texttt{xvar()} is age, L, M, and S are tabulated at monthly intervals. For height and length, these parameters are tabulated at 1 cm intervals. Where a child’s age/height/length occurs within these intervals, values of L, M, and S are obtained via a linear interpolation between successive values of \texttt{xvar()}. (The BMI cutoff points are tabulated at 6 monthly intervals. The cutoff point where a child’s age occurs within these intervals is also obtained via linear interpolation.) Where \texttt{xvar()} is outside the permitted range, \texttt{zanthro} and \texttt{zbmicat} return a missing value.

In the British charts, the L value was fixed at 1 for height, head circumference, sitting height, and leg length, since these measures are normally distributed.
Table 1: 2000 CDC Growth Charts: Version US

<table>
<thead>
<tr>
<th>Chart code</th>
<th>Description</th>
<th>Measurement unit</th>
<th>xvar() range</th>
</tr>
</thead>
<tbody>
<tr>
<td>ha</td>
<td>height-for-age</td>
<td>cm</td>
<td>2 to 20 years</td>
</tr>
<tr>
<td>wa</td>
<td>weight-for-age</td>
<td>kg</td>
<td>0 to 20 years</td>
</tr>
<tr>
<td>ba</td>
<td>bmi-for-age</td>
<td>kg/m²</td>
<td>2 to 20 years</td>
</tr>
<tr>
<td>hca</td>
<td>head circumference-for-age</td>
<td>cm</td>
<td>0 to 36 months</td>
</tr>
<tr>
<td>la</td>
<td>length-for-age</td>
<td>cm</td>
<td>0 to 36.5 months</td>
</tr>
<tr>
<td>wh</td>
<td>weight-for-height</td>
<td>kg</td>
<td>77 to 121.5 cm</td>
</tr>
<tr>
<td>wl</td>
<td>weight-for-length</td>
<td>kg</td>
<td>45 to 103.5 cm</td>
</tr>
</tbody>
</table>

Table 2: 1990 British Growth Charts: Version UK

<table>
<thead>
<tr>
<th>Chart code</th>
<th>Description</th>
<th>Measurement unit</th>
<th>xvar() range</th>
</tr>
</thead>
<tbody>
<tr>
<td>ha</td>
<td>height-for-age</td>
<td>cm</td>
<td>0 to 23 years</td>
</tr>
<tr>
<td>wa</td>
<td>weight-for-age</td>
<td>kg</td>
<td>0 to 23 years</td>
</tr>
<tr>
<td>ba</td>
<td>bmi-for-age</td>
<td>kg/m²</td>
<td>0 to 23 years</td>
</tr>
<tr>
<td>hca</td>
<td>head circumference-for-age</td>
<td>cm</td>
<td>Males: 0 to 18 years</td>
</tr>
<tr>
<td>sha</td>
<td>sitting height-for-age</td>
<td>cm</td>
<td>Females: 0 to 17 years</td>
</tr>
<tr>
<td>lla</td>
<td>leg length-for-age</td>
<td>cm</td>
<td>0 to 23 years</td>
</tr>
</tbody>
</table>

6 Examples

We illustrate with some data on a set of Australian children aged 5 to 10 years. Below, we show the relevant data for the first 5 children. The variable `sex` is coded male = 1, female = 2.

```
.list in 1/5, noobs
```

```
<table>
<thead>
<tr>
<th>id</th>
<th>sex</th>
<th>ageyrs</th>
<th>bmi</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>6.6</td>
<td>15.5</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>8.4</td>
<td>20.1</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>5.7</td>
<td>16.8</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
<td>7.5</td>
<td>19.9</td>
</tr>
<tr>
<td>5</td>
<td>2</td>
<td>9.7</td>
<td>26.8</td>
</tr>
</tbody>
</table>
```

In order to compare the BMI of the children in this sample, for instance, with respect to socioeconomic grouping or between two time points several years apart, the BMI values may be converted to standardized z-scores. The z-scores are created using the following command:
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```
. egen zbmiuk = zanthro(bmi, ba, UK), xvar(ageyrs) gender(sex)
    gencode(male=1, female=2)

For this analysis of Australian children, it might be equally appropriate to use the US reference standards:

. egen zbmius = zanthro(bmi, ba, US), xvar(ageyrs) gender(sex)
    gencode(male=1, female=2)

To determine the proportion of children who are overweight or obese, each child can be categorized using the following command:

. egen bmicat = zbmicat(bmi), xvar(ageyrs) gender(sex)
    gencode(male=1, female=2)

Here are the results for the first 5 children:

. list in 1/5, noobs

<table>
<thead>
<tr>
<th>id</th>
<th>sex</th>
<th>ageyrs</th>
<th>bmi</th>
<th>zbmiuk</th>
<th>zbmius</th>
<th>bmicat</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>6.6</td>
<td>15.5</td>
<td>-.0172949</td>
<td>.0460011</td>
<td>Normal wt</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>8.4</td>
<td>20.1</td>
<td>1.921224</td>
<td>1.573253</td>
<td>Overweight</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>5.7</td>
<td>16.8</td>
<td>.8006477</td>
<td>.9584172</td>
<td>Normal wt</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
<td>7.5</td>
<td>19.9</td>
<td>2.08305</td>
<td>1.729176</td>
<td>Overweight</td>
</tr>
<tr>
<td>5</td>
<td>2</td>
<td>9.7</td>
<td>26.8</td>
<td>2.803655</td>
<td>2.183658</td>
<td>Obese</td>
</tr>
</tbody>
</table>
```

An example where the age variable is coded in months and the outcome of interest is weight would be handled as follows:

```
. egen zwauk = zanthro(weight, wa, UK), xvar(agemth) gender(sex)
    gencode(male=1, female=2) ageunit(month)

Using the US standards for height or length:

. egen zwhus = zanthro(weight, wh, US), xvar(height) gender(sex)
    gencode(male=1, female=2)
. egen zulus = zanthro(weight, wl, US), xvar(length) gender(sex)
    gencode(male=1, female=2)
```

With the variable sex coded male = M, female = F:

```
. egen zbauk1 = zanthro(bmi, ba, UK), xvar(agemth) gender(sex)
    gencode(male=M, female=F) ageunit(month)
. egen bcat = zbmicat(bmi), xvar(agemth) gender(sex)
    gencode(male=M, female=F) ageunit(month)
```

Now, swapping the order in specifying the gender codes and omitting the comma:

```
. egen zbauk1 = zanthro(bmi, ba, UK), xvar(agemth) gender(sex)
    gencode(female=F male=M) ageunit(month)
```
and using just the first letter in specifying each gender:

```
. egen bcat = zbmicat(bmi), xvar(agemth) gender(sex)
  > gencode(f=F m=M) ageunit(month)
```

NOTE: The code cannot be abbreviated. It must be typed into the `gencode()` option exactly as it appears in your dataset; e.g., `sex` is coded male = Male, female = Female:

```
. egen bcat = zbmicat(bmi), xvar(agemth) gender(sex)
  > gencode(f=Female m=Male) ageunit(month)
```

and forcing calculation of all $z$-scores:

```
. egen zbauk2 = zanthro(bmi,ba,UK), xvar(agemth) gender(sex)
  > gencode(f=Female m=Male) ageunit(month) nocutoff
```

7 References


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