Estimation of Adherence to Antipsychotic and Diabetic Medications in a Sample of Schizophrenia Patients

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ABSTRACT

We studied a sample of Schizophrenia patients and were interested in knowing (1) Are schizophrenia patients adherent to their medication if they are also taking diabetic medication? And (2) How adherent to their medication are Schizophrenia patients who have no comorbidity and are their significant differences between these two groups? (3) Can we work with a simplified data structure using multiple records per patient and attain efficiency while maintaining accuracy?

The results indicate that Schizophrenia patients with no comorbidity are less likely to be adherent to their medication than patients on multiple medications. However, this difference was not statistically significant.

We were able to work with a simplified data structure and were able to achieve efficiency in calculations using advanced techniques to calculate the medication possession ratio (MPR).

INTRODUCTION

Non-adherence to medication is a significant health care cost. Roughly a quarter of the patients on medication are non-compliant (Bovet et al., 2002). The New England Health Care Institute (NEHI) has recently estimated the annual cost of medication non-adherence at $290 Billion (August 11, 2009). The non-adherence issue is recognized as a significant problem by the International Society for Pharmacoeconomics and Outcomes Research (ISPOR). It has set up special task groups who have initiated studies to seek a greater understanding of the problem.

BRIEF REVIEW OF THE LITERATURE

While there are extensive studies on the medication adherence problem and the economic costs associated with it, the method of estimation has received limited coverage in the SAS literature. Estimating medication adherence is a complicated task. Sophisticated algorithms have been developed to capture the Medication Possession Ratio (MPR) which is a measure of medication adherence (Leslie, 2007). However, medication adherence has been primarily examined from the point of view of a single drug. What happens when the patient is on multiple drugs?

There is recent evidence that patients on single medication for a chronic condition are less adhering to medication than those on multiple medications (Kreyenbuhl et al., 2010).

We will briefly touch upon the study design, methods and results which may be published in detail elsewhere later. The main focus of this paper will be on the SAS® use for our study. We present a different approach than using array manipulation for Medical Possession Ratio calculations. We explain the merits of using a narrow and long dataset as opposed to a wide data set typically used for extensive array manipulation. We also briefly demonstrate the efficiencies gained using the hash data step object for looking-up subsets. Finally, we present results using re-sampling of the data to guard against spurious results (Wesfall, 1999).

STUDY DESIGN

The MarketScan® Commercial Claims & Encounters data for years 2006 and 2007 were restricted to a sample of Schizophrenia patients, <65 years of age, who were also on medication for diabetes control. The data were cleaned for quality control.

We hypothesize based on the recent study that Schizophrenia patients on single medication are less likely to be adherent to their medication than those with co-morbidity and on multiple medications. Therefore, we expect to see lower adherence among Schizophrenia patients who were not on any other another chronic medication (n=444) compared to those who were also on Diabetic control medication (n=864). We define not being on any another chronic medication as follows. The number of days in possession of any non-antipsychotic medication must not exceed one quarter of the number of days in possession of an antipsychotic medication in a year.
SELECTION OF PATIENTS

We filtered the patients in the following order.

1. Schizophrenia cases: patients with at least one inpatient or outpatient medical claim with a diagnosis of schizophrenia (ICD-9 codes 295.xx) and at least one prescription for an antipsychotic medication in 2006. Cases were further restricted to those individuals participating in health plans that reported prescription drug data for both 2006 and 2007.

2. Subset who were on Diabetic medication (had at least one prescription for a diabetes medication in 2006)

3. Subset who were NOT on Diabetic medication (mutually exclusive from 2)

4. Subset Diabetic patients to those who also took Antipsychotic medication.

5. Subset NON-Diabetic patients to those who took Antipsychotic medication and no other chronic medication.

Since these are large data sets we had to be mindful of efficient selection processes. Given below are the salient parts of the code with brief description headers.

```sql
%*-------------------------------------------------- -----------------;
%*   Assign values to a macro variable for repeated KEEP statements  ;
%let keeplist = enrolid svcdate ndcnum daysupp thercls ;
%*-------------------------------------------------- -----------------;
%*   Extract Schizophrenia patients taking diabetic medication       ;
%*-------------------------------------------------- -----------------;
data diablist ;
  set schizo.ccaed062_schizo (keep=&keeplist)  
    schizo.ccaed071_schizo (keep=&keeplist)  ;
  by enrolid ;
  where not missing(enrolid) and  
    sign(daysupp) >-1 and 
    thercls in (172,173,174) ;
run;
%*-------------------------------------------------- -----------------;
%*   Get a list of the IDs to load to a memory table to extract      ;
%*   subsets.                                                        ;
%*-------------------------------------------------- -----------------;
%proc sql nobprint ;
  create table diabidlist as 
    select distinct enrolid  
      from diablist ;
  select count(distinct enrolid)  
    into : idcount separated by " "  
      from diablist ;
quit ;
%put <&idcount> ;
%*-------------------------------------------------- -----------------;
%*   Extract patients taking antipsychotic medications               ;
%*   limiting them to the subset taking diabetic medication.         ;
%*   The Data Step Hash Object is a good candidate for efficiency    ;
%*   Define the Hash objects on the first iteration of the datastep. ;
%*-------------------------------------------------- -----------------;
```
data psychanddiab ;
  if _N_ = 1 then do;
    declare Hash ht(dataset: "diabidlist"); -----------------------------(1)
    ht.defineKey("enrolid"); ----------------------------------------(2)
    ht.defineData("enrolid"); ---------------------------------------(3)
    ht.defineDone(); -------------------------------------------------(4)
  end;
/* Load enrolid from both and query hash table */
/* This should give all patients who took THERCLS=70 */
set schizo.ccaed062_schizo (keep=&keeplist)
  schizo.ccaed071_schizo (keep=&keeplist) ;
by enrolid ;
where not missing(enrolid) and
  sign(daysupp) >=-1 and
  thercls = 70 ;
if ht.find() = 0; *patients in IDLIST; --------------------------(5)
run;

A brief explanation about the Hash object used above. This is a relatively new feature in SAS introduced in Version 9. When merging two datasets, if one dataset is small compared to other, the entire small dataset can be loaded to memory and looked up by the same key of the larger dataset. This makes the process much quicker. There were a number of papers using the hashing method with temporary arrays even before the introduction of the hash object (e.g. Dorfman, 2001). In the code above the reference line (1) declares the DIABIDLIST dataset as the one to load in memory. The key on which the larger dataset(s) will search is declared in (2) as ENROLID. The same variable is declared as satellite data to be carried with it to the output dataset PSYCHANDDIAB. Finally (4) says we have done defining our hash table. Now the job of the larger dataset is to go find a match on ENROLID which is stated in (5). When the ID is found a return code of 0 is returned indicating success and the record is output to the dataset PSYCHANDDIAB. There are many complex pieces to the Hash object and the reader is referred to a nice introduction by Secosky and Bloom (2006).

SELECTION OF COMPARISON PATIENTS

The Schizophrenia patients with diabetes were compared to Schizophrenia patients without diabetes or any other chronic medication use. We had operationally defined this as no other medication should be filled more than 25 percent of the days that antipsychotic medications were filled for that patient. If exceeded that patient is dropped. So this could be a tricky selection but could be simplified keeping the same data structure with multiple rows per patient. However, a temporary array is used to load the days supplied (DAYSUPP) that was summed by therapeutic class (THERCLS) and given a new name SUMDAYSUPP. Our reference SUMDAYSUPP is therapeutic class value 70 for antipsychotic medication. Once the data step reads all the records for the patient, we examine every element of the array at the end of the patient’s record. Please note that this type of DO loop with a SET inside it is not conventional use. However, we can take advantage of certain desirable properties to limit the number of steps in the coding. For a more complete discussion of this DO loop structure, see Chakravarthy (2003) and Dorfman and Vyverman (2009).

%*-------------------------------------------------------------;
%*   Sum up the days supplied by enrolid by thercls. This allows  
%*   us to examine later for other medications for chronic use.   
%*-------------------------------------------------------------;
var array lug (300) _temporary_; 
/* load all DAYSUPP by enrolid */
do until (last.enrolid) ;
  set restpsych ;
  by enrolid thercls ;
  lug(thercls) = sumdaysupp ;
if thercls = 70 then reference = sumdaysupp;
end;

/* initialize for selection by default and mark for deletion only when the deletion criteria is met array element with index value 70 is antipsychotic which is the reference value. This should not be evaluated */
todelete = 0;
do _n_ = 1 to dim(lup);
if not todelete then do;
    if not missing(lup(_n_)) and _n_ ^= 70 and (lup(_n_) > (reference) or lup(_n_) > (.25*reference)) then do;
        todelete = 1;
tocheckdaysupp = lup(_n_);
tochecktherls = _n_;
    end;
end;
lup(_n_) = .;
end;
run;

**CALCULATION OF THE MEDICATION POSSESSION RATIO (MPR)**
Once the assembling of the data was completed, it was time to calculate the medication possession ratio (MPR). We used the ISPOR definition:

Number of Days of Medication Supplied within the Refill Interval/Number of Days in Refill Interval.

In past SAS papers, the use of ARRAYS was a popular choice for this calculation. In this study we have used logic similar to that but have worked with the simpler data structure of multiple records per patient. This is easier to view when printed or seen through a VIEWTABLE window. ARRAYS are still used but only for populating the calculated MPR values to the different Medications. Here is the single datastep that calculates MPR.

data mpr5;
do until (last.name);
    set temp5_3;
    by enrolid name svcdate;
    /* once the refill date goes past a year from first prescription we want to stop processing. The CONTINUE statement continues with the DO loop without doing anything further.*/
    if stopProcessDate then continue;
    if first.name then do;
        cumdays = daysupp;
totdays = daysupp;
        firstSupplyDate = svcdate;
        /* This is the cutoff date */
        AnniversaryDate = (firstSupplyDate + 365) - 1;
        end;
    if svcdate >= AnniversaryDate
then stopProcessDate = svcdate;
if stopProcessDate then continue;

if not first.name and
   not (first.name and last.name) and
   (svcdate - PrevServiceDate > (.5*PrevSupply))
   then cumdays = sum(cumdays, daysupp);
   PrevServiceDate = svcdate;
   PrevSupply = daysupp;
end;

array allrx(*) &druglist;
array psych(*) &druglist2;
retain allrx psych;

if stopProcessDate
   then cumdays = cumdays - prevsupply +
       (AnniversaryDate-PrevServiceDate+1);

if cumdays then delete;
allrx(input(name, indrug.)) = cumdays / 365;
if name in (&psychdrugs) then
   psych(input(trim(name)||"2",indrg.)) = cumdays;

/* This calculates the MPR for all of ATYPICAL and non-ATYPICAL
   antipsychotics combined */
if last.enrolid then
   do;
      psychMPR = (sum(of psych(*))) / 
                  ((dim(psych)-nmiss(of psych(*))) * 365);
      output;
      do _n_ = 1 to dim(allrx);
         allrx(_n_) = .;
      end;
      do _n_ = 1 to dim(psych);
         psych(_n_) = .;
      end;
   end;
   format AnniversaryDate PrevServiceDate firstSupplyDate
      stopProcessDate mmddyy10.
   keep enrolid ndcnum svcdate daysupp prodnme name
      AnniversaryDate PrevServiceDate firstSupplyDate
      stopProcessDate cumdays &druglist &druglist2 psychMPR
      PrevSupply;
run;

METHODS AND DISCUSSION OF RESULTS
The descriptive statistics of the demographics in the study is produced below in tables 1 and 2.
**Table 1: Descriptive Statistics on Age Distribution**

<table>
<thead>
<tr>
<th>Age of Patient</th>
<th>N</th>
<th>Mean</th>
<th>Std Dev</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1308</td>
<td>45.1437</td>
<td>12.887</td>
<td>10.000</td>
<td>63.000</td>
</tr>
</tbody>
</table>

**Table 2: Frequency Distribution of Gender**

<table>
<thead>
<tr>
<th>Gender of Patient</th>
<th>SEX</th>
<th>Frequency</th>
<th>Percent</th>
<th>Cumulative Frequency</th>
<th>Cumulative Percent</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1- M</td>
<td>606</td>
<td>46.33</td>
<td>606</td>
<td>46.33</td>
</tr>
<tr>
<td></td>
<td>2- F</td>
<td>702</td>
<td>53.67</td>
<td>1308</td>
<td>100.00</td>
</tr>
</tbody>
</table>

The MPR ratios were calculated individually for each Atypical and Non-Atypical medication. The oral diabetic medications were combined and the Insulin class was separated. The mean Medication Possession Ratio is displayed in Table 3.

**Table 3: Mean Medication Possession Ratios**

<table>
<thead>
<tr>
<th>Analysis Variable : Medication Possession Ratio</th>
<th>Name of Medication</th>
<th>N Obs</th>
<th>Lower 95% CL for Mean</th>
<th>Upper 95% CL for Mean</th>
<th>Mean</th>
<th>Std Dev</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ABILI</td>
<td>266</td>
<td>0.5360</td>
<td>0.6194</td>
<td>0.5777</td>
<td>0.3453</td>
</tr>
<tr>
<td></td>
<td>CLOZA</td>
<td>87</td>
<td>0.7036</td>
<td>0.8304</td>
<td>0.7670</td>
<td>0.2974</td>
</tr>
<tr>
<td></td>
<td>GEODO</td>
<td>148</td>
<td>0.5228</td>
<td>0.6398</td>
<td>0.5813</td>
<td>0.3601</td>
</tr>
<tr>
<td></td>
<td>INVEG</td>
<td>27</td>
<td>0.1657</td>
<td>0.3167</td>
<td>0.2412</td>
<td>0.1909</td>
</tr>
<tr>
<td></td>
<td>RISPE</td>
<td>282</td>
<td>0.6021</td>
<td>0.6793</td>
<td>0.6407</td>
<td>0.3293</td>
</tr>
<tr>
<td></td>
<td>SEROQ</td>
<td>275</td>
<td>0.5749</td>
<td>0.6564</td>
<td>0.6156</td>
<td>0.3433</td>
</tr>
<tr>
<td></td>
<td>ZYPRE</td>
<td>143</td>
<td>0.5594</td>
<td>0.6709</td>
<td>0.6151</td>
<td>0.3373</td>
</tr>
<tr>
<td></td>
<td>OTHER</td>
<td>288</td>
<td>0.5683</td>
<td>0.6473</td>
<td>0.6078</td>
<td>0.3407</td>
</tr>
<tr>
<td></td>
<td>INSULIN</td>
<td>238</td>
<td>0.5198</td>
<td>0.5980</td>
<td>0.5589</td>
<td>0.3059</td>
</tr>
<tr>
<td></td>
<td>ORALDIAB</td>
<td>795</td>
<td>0.6905</td>
<td>0.7322</td>
<td>0.7114</td>
<td>0.2996</td>
</tr>
<tr>
<td></td>
<td>All Antipsychotics</td>
<td>864</td>
<td>0.6501</td>
<td>0.6881</td>
<td>0.6691</td>
<td>0.2845</td>
</tr>
<tr>
<td></td>
<td>Antipsychotics No Co-morbidity</td>
<td>444</td>
<td>0.5899</td>
<td>0.6491</td>
<td>0.6195</td>
<td>0.3173</td>
</tr>
</tbody>
</table>

We were interested in knowing how adherent are Schizophrenia patients with (a) their antipsychotic medications and (b) their diabetic medication. We were also interested in knowing how adherent to medications are Schizophrenia patients having no other co-morbidity. Based on a recent publication, patients on single medication are less likely to be adherent than those using multiple drugs. Preliminary indication points to the same. Patients on single medication appear to be less likely to be adherent to their medication than patients with co-morbidity and on multiple chronic medications. However, this needs to be tested for statistical significance. We follow Pearce and Wesfall (1998) recommendation and use PROC MULTTEST resampling without replacement.
Table 4: Test of Differences in Mean Medication Possession Ratios

<table>
<thead>
<tr>
<th>Variable</th>
<th>Contrast</th>
<th>Raw</th>
<th>Stepdown Permutation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medication Possession Ratio</td>
<td>OralDiab vs Psych</td>
<td>0.0069</td>
<td>0.0137</td>
</tr>
<tr>
<td></td>
<td>OralDiab vs PsychONLY</td>
<td>&lt;.0001</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td></td>
<td>Psych vs PsychONLY</td>
<td>0.1280</td>
<td>0.1278</td>
</tr>
</tbody>
</table>

Even though preliminary indication was that schizophrenic patients with no co-morbidity were less likely to be adherent to their medications than patients on multiple chronic medications, the difference is not statistically significant. It is to be noted that a test of the mean differences using the same MULTTEST procedure yielded a statistically significant differences between the two groups. However, we followed Pearce and Westfall’s recommendation to use the Cochran-Armitage linear trend tests for group comparisons. Patients in the study were more likely to be adherent to their diabetic medication than their antipsychotic medication. This was statistically significant.

CONCLUSION

We had hypothesized that Schizophrenia patients with no other co-morbidity are less likely to adhere to their medication than patients who were on multiple medications. This logic was based on the results from a recent publication. Although this appeared true in our study the difference was not statistically significant.

The main focus of this paper was to demonstrate efficiencies attained in using certain SAS code. We also demonstrated a different method of calculating MPR using the standard definition. In the process we showed how a narrow and long data structure is more easy to handle than a wide and short data structure.

REFERENCES


Pearce, Gregory L. and Westfall, Peter H. (1998). Using Resampling Techniques in PROC MULTTEST to Evaluate Surgeon Specific Results Following Coronary Artery Bypass Graft (CABG) Surgery

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