Environmental Factors Effects on Asthma in the National Population Health Survey

Environmental Factors Effects on Asthma in the National Population Health Survey

By

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A Project

Submitted to the School of Graduate Studies

In Partial Fulfillment of the Requirements

for the Degree

Master of Science

McMaster University

March, 2003

MASTER OF SCIENCE (2003) (Statistics)

McMaster University Hamilton Ontario

TITLE:Environmental Factors Effects on Asthma
in the National Population Health Survey

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NUMBER OF PAGES: x, 104

Abstract

Asthma is one of the most common respiratory diseases in Canada. It not only brings pain to more than 7.8% of Canadians but it also costs millions of dollars every year.

In this project we study the influence that environmental factors have on asthma based on the data from the National Population Health Survey (NPHS) conducted by Statistics Canada. A descriptive analysis is done first to get an initial understanding of the environmental factors' effects on asthma. Then we do a χ^2 -test to test for the homogeneity of asthma distribution across the levels of each environmental factor. Most of the factors included in our study are significant except those representing whether living in the metropolitan areas of Montreal or Vancouver, the number of persons living in the household, and whether there are small children in the household.

For ordinal variables we test for trend on asthma prevalence. The trend tests indicate that there are significant trends between asthma and most of the ordinal factors except for a few, including the number of bedrooms in the household and the number of cigarettes smoked daily by the daily smokers.

Then odds ratio and relative risk analyses are done to obtain statistical insights on the relative risk of the factors. The result shows that living in Nova Scotia, Ontario and Quebec, urban areas, engaged in finance, community services, personal service, young, attend physical activities, born in Canada, white, single or widowed, separated, divorced, start smoking early, do not own the dwelling living in, female adult, male children, and overweight and underweight adult, are all contributing factors for asthma.

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To model and investigate the joint effect of factors on asthma prevalence, we use logistic and log-linear regression models. To avoid collinearity problems, a reduced number of predictors is used. The results suggest that environmental factors have a significant joint influence on asthma prevalence.

Acknowledgements

I am very grateful to have this opportunity to express my deep appreciation to my supervisor, Dr. Román Viveros-Aguilera, for his inspiring direction and support throughout the entire process of this project. Dr. Román Viveros-Aguilera is a patient and understanding supervisor. I am very fortunate to have his great guidance.

Many thanks to Mr. Vivek Jadon from the Data/Text Center at Mills Library, McMaster University for his enthusiasm and help during the initial stages of my project.

I thank Dr. Peter Macdonald and Dr. Lehana Thabane for their careful reading of my project and for their insightful suggestions that led to many improvements in the final version of my project.

I am also grateful to my professors, staff and my fellow graduate students in the Department of Mathematics and Statistics of McMaster University. It was them who made my studies enjoyable and worthwhile.

I am extremely thankful to my parents for their encouragement and support. A special thanks to my husband for his deep love.

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Chapter 1: The National Population Health Survey (NPHS): Data and Study Objectives

1.1 Introduction

The National Population Health Survey (NPHS) is designed to collect information related to the health of the Canadian population. The survey began in 1994, and conducted and managed by Statistics Canada by-yearly. It is composed of three component parts: the survey of households, the survey of institutions and the survey of the North (Yukon and the Northwest Territories with the principal exclusion of populations on Indian Reserves, Canadian Forces Bases and some of the most northerly remote areas of the Territories.). The household component includes household residents in all provinces, with the exclusion of populations on Indian Reserves, Canadian Forces Bases and some remote areas in Québec and Ontario. In each household, some limited information was collected from all household members (general component) and one person in each household was randomly selected for a more in-depth interview (health component). The 1996-97 NPHS was collected mainly by telephone. For that reason, an effort has been made to make the questions easier to read (for the interviewers) and easier to understand (for the respondents) by using more colloquial wording where possible. Without revamping sections, screens have been redesigned, response categories have

been shortened and frequent responses in "other specify" have become itemized categories.

The source data for our study come from the 1996-97 NPHS Public Use Microdata Documentation. It should be noted that the "public use" microdata files differ in a number of important respects from the survey "master" files held by Statistics Canada. These differences are the result of actions taken to protect the anonymity of individual survey respondents through suppression of individual values, variable grouping, and variable capping.

The principle behind estimation in a probability sample such as the NPHS is that each person in the sample has a sampling weight. WT66 is the principal weight variable on the Health File and should be used for analysis of most variables. For many analysis techniques (for example linear regression, logistic regression, analysis of variance), a method exists that can make the application of standard packages more meaningful. If the weights on the records are rescaled so that the average weight is 1, then the results produced by the standard packages will be more reasonable since they take into account the unequal probabilities of selection. The rescaling can be accomplished by using in the analysis a weight equal to the original weight divided by the average of the original weights for the sampled units (people) contributing to the estimator in question. This approach was taken in this project.

As indicated earlier, the data came from Statistics Canada, one of the most respectable institutions in the country for conducting surveys. Since I did not have access to the original surveys, it was not possible to verify the reliability of the data.

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1.2 Case and Variable Selection, Sub-Data Description

The NPHS data are stored into two different data sets. The information collected about all household members is stored on the General File. Generally, in each household, one person was selected to answer a more in-depth questionnaire related to health. These data are stored on the Health File.

Each record on the Health File corresponds to the household of the selected respondent. This file carries all data collected pertaining to the selected respondent, e.g. all the variables from the General File belonging to the selected respondent and all variables resulting from the in-depth questionnaire, such as smoking, stress, mental health, etc. Since the Health File contains more information, we chose it for our main study.

The data analyzed in this project are the 1996-1997 NHPS data obtained in CDs from the Data/Text Center, Mills Library, McMaster University. The main file is the Health File which is stored as a text file named H356.txt. Using the SAS program listed in the Appendix B we unzipped and restored the data set. The Health File data contains 81,804 records and 944 variables. The 944 variables relate to the following characteristics: alcohol dependence, attitudes towards parents, blood pressure, chronic conditions, coping, drug use, demographic and household variables, dental visits, education, emergency services, eye examination, flu shots, geographic identifiers, general health, health care utilization, health information, health status, HIV, height and weight, injury, income, insurance, labor force, mental health, physical activity, physical check-up, province, restriction of activity, repetitive strain, road safety, socio-demographics, sexual health, smoking, social support, health services, training and UV exposure, two-week disability, violence and personal safety, women's health, and sample weights.

Because of its known high prevalence, particularly in children, asthma has been an intensively studied disease. Rees and Kanabar (2000) report that ethnicity, family history, rural / urban environment, body weight, indoor environment (particularly for children), vigorous exercise, occupation, emotional factors, pollution, weather and air quality all have an influence on asthma. The studies reported vary in their focus of analysis and methods, and usually based on data from specific parts of the world. Manning (1993) discuses the effects that different forms of exercise have on asthma. One of his conclusions is that more than 50% of asthmatic subjects experience progressively less and less airway narrowing with repeated exercise challenges on the same day. Exercise induced asthma is also studied by Tan and Spector (2002). Other studies focus on the effect on asthma of more restricted environments such as school (Leickly, 2003) and urban/rural areas (Maffei et al., 2001). Since our main objective is to study the environmental effects on asthma prevalence, we need not to include all the 944 variables from the survey. After reviewing the survey questionnaire, the meaning of each variable and our study objectives, 35 variables are selected for study. So a sub data set from the main source data (Health File) is used in our study to do the analysis. The sub data still has 81,804 records but with only the 35 variables which are of interest in our study. The variables included in our sub data can be divided into groups that describe outdoor environment, indoor environment, working environment, physical exercise and air exchange amount, social demographics and whether in a smoking environment. We also include variables that describe basic characteristics of respondents in our sub data.

The selected variables are listed in Table 1.1. For details about variable meaning, coding,

range and universe please see Appendix A.

\sim	<u> </u>	
	Variable Code	Description
Main dependent variable	CCC6_1C	Has asthma (Do you have asthma diagnosed by a health professional? Yes or No)
	PRC6_CUR	Province of residence
Outdoor environment	GE36GCMA	Derived 1991 Census Metropolitan Area - grouped
variables	GE36GHLR	Derived health areas – grouped (26 groups)
	GE36GHRO	Derived health areas – grouped (33 groups)
	GE36GURB	Derived rural and urban area – grouped
Working environment	LFC6GO21	Occupation Codes for main job – grouped (21 groups)
variables	LFC6GI13	Industry Codes for main job – grouped (13 groups)
	PAC6DEE	Derived energy expenditure (1 decimal point)
Physical exercise, air	PAC6DLEI	Derived participant in leisure physical activity
exchange amount	PAC6DFM	Derived monthly freq. of physical activity lasting >15min.
(respiratory system)	PAC6DFR	Derived frequency of all physical activity
	PAC6DFD	Derived participation in daily phys. Activities > 15 min.
	PAC6DPAI	Derived physical activity index
	SDC6GCB	Country of birth – grouped
Demographic variables	SDC6GRAC	Derived race or color – grouped
	DHC6GMAR	Marital status – grouped

 Table 1.1
 List of variable names and meaning descriptions.

	Variable Code	Description
Whether in smoking	SMC6_3	Age started smoking daily – daily smoker
environment	SMC6_4	Number of cigarettes smoked each day – daily smoker
	DHC6_OWN	Dwelling owned by household member
	DHC6GBED	Derived number of bedrooms in dwelling – grouped
Indoor environment	DHC6GHSZ	Derived household size – grouped
variables	DHC6GLE5	Derived persons <= 5 years old in household – grouped
	DHC6G611	Derived persons 6 to 11 years old in hhld – grouped
	DHC6GECF	Derived household type – grouped
	DHC6DLVG	Derived living arrangements of the selected respondent
	HWC6GHT	Height - adults and children - grouped
	HWC6GSW	Derived standard weight - grouped
Basic variables of	HWC6G3KG	Weight in kilograms - grouped
respondents'	HWC6GBMI	Derived Body Mass Index (1 decimal place) - grouped
characteristics	DHC6_SEX	Gender
	DHC6GAGE	Age - grouped
	CCC6_1H	Has chronic bronchitis or emphysema
	DGK6_1	Takes ventolin or other inhalants
Sampling weight	WT66	Sampling weight for selected respondent

1.3 Study Objectives

The overall objectives of this project are to study the influence of environmental factors on asthma. More concretely, our main aims can be described as follows.

- (a) To assess individually which of the factors studied have a significant influence on asthma prevalence. This will be accomplished by using appropriate graphical methods and by a Chi-square test.
- (b) Detect trends between asthma prevalence and ordinal categorical and non categorical factors. The Cochran-Armitage test will be used.
- (c) For factors that have a significant influence on asthma prevalence from (a), study the relevant odds ratios and relative risks by constructing 2×2 tables.
- (d) Relate the factors in a joint manner to the asthma prevalence through logistic regression models and do the associated estimation and inference. First, the model will be applied to the whole data. Then the data will be split into infants, children and adults. Each of these groups will be randomly partitioned into three subsets, one for variable selection, the second one for model fitting and the third are for cross-validation.

1.4 Descriptive Analysis of Data

1.4.1 Response Variable CCC6_1C

Table 1.2 displays the sample frequency distribution for the response variable in this project, CCC6_1C (Has Asthma). The corresponding general population frequency

distribution data are from the 1996 national census of Canada. All information in Table 1.2 can be found in the data dictionary of 1996-1997 NPHS survey. Most respondents have unambiguous answers on whether they have asthma or not. The respondents who answer "Don't know", "Refusal" and "Not stated" account for only 50 out of 81,804, that is, 0.061% of all the respondents. To simplify the analysis we will delete those respondents who do not answer "Yes" or "No" on whether they have asthma. So in the sub data we study, there are 81754 cases. Note that among the Yes / No respondents, 7.64% and 7.81% have asthma in the sample and in the population, respectively.

Variable Name	CCC6_1C	Length	1	Position	165
Question Name	CHR-Q1				
Concept	Has asthma				
Duestion	Do you have asthma	diagnosed by a health pr	ofessional?		
Universe	All respondents				
Note					
<u>Content</u>			Code	Sample	Population
YES			1	6,242	2,236,139
NO			2	75,512	26,394,976
DON'T KNOW			7	38	7,859
REFUSAL			8	5	523
NOT STATED			9	7	2,239
		Total		81,804	28,641,736

 Table 1.2 Frequency distribution for variable CCC6_1C (Has Asthma).

1.4.2 Independent Variables

In the following we review the sub data from the seven groups of variables described above. Most of the graphs were generated using Splus and Excel.

The NPHS survey shows that the overall asthma prevalence rate in Canada is 7.81%. Table 1.3 depicts the asthma rate per province while Figure 1.1 displays the corresponding histogram. However, the prevalence rate distribution among provinces and health areas is different. The highest occurs at the province of Nova Scotia, which is as

high as 8.43%. The second highest is Ontario, which is 8.04%. The lowest ones are Newfoundland and New Brunswick, with 5.76% and 6.6%, respectively.

Province	Asthma Prevalence (%)	Percentage of people with asthma in each province to the total number of asthma suffers from all provinces (%)
Nova Scotia	8.43	3.38
Ontario	8.04	38.95
Quebec	7.98	25.16
P.E.I.	7.78	0.46
British Columbia	7.76	12.79
Alberta	7.52	9.17
Manitoba	7.39	3.58
Saskatchewan	6.95	2.95
New Brunswick	6.6	2.15
Newfoundland	5.76	1.41
Whole Canada	7.81	100

 Table 1.3 Asthma rate per province.

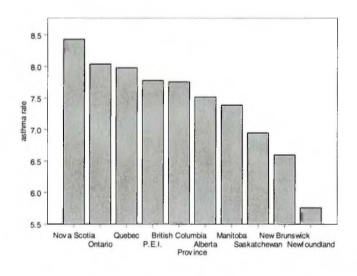


Figure 1.1 Asthma rate for the Canadian provinces.

Since the provinces vary dramatically in size, so do the total count of asthma sufferers. These variations, which appear in column 3 of Table 1.3 are plotted in Figure 1.2. Note these figures are the gross percentages, they have not been adjusted for provinces size.

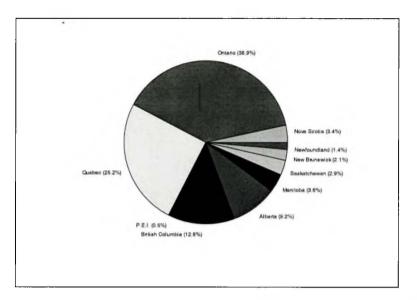


Figure 1.2 Pie chart of the percentage of asthma sufferers per province.

Variable GE36GURB (whether living in urban or rural area) provides another view of asthma distribution. People living in urban areas have an asthma rate of 7.88%, whereas people living in rural areas have the smaller rate of 6.52%. This partly means that a rural environment is a little "healthier" than an urban environment for asthma sufferers.

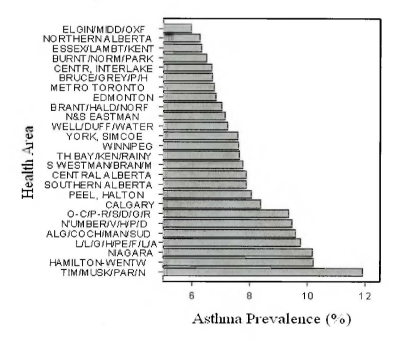


Figure 1.3 Asthma rate among health areas for Ontario, Manitoba and Alberta.

Some provinces such as Ontario, Manitoba and Alberta provide larger financial and human resources thus more detailed surveying is done. The three provinces account for more than a half of the total population. The three provinces are divided into 26 health areas; the histogram in Figure 1.3 displays the asthma rates among the 26 areas.

It is clear that the asthma rate is not uniform across areas. From the frequency table we note that the asthma rate is as low as 5.96% for the area of ELGIN, MIDDLESEX and OXFORD and as high as 11.93% for the area of TIMISKAM., MUSKOKA, PARRY SOUND and NIPISS.

Occupational asthma is a disease characterized by variable air flow limitation or airway hyper-responsiveness due to causes and conditions attributable to a particular occupational environment and not to stimuli encountered outside the workplace (Bernstein *et al.* 1993).

Variable LFC6GO21 has 21 groups (Figure 1.4) and LFC6GI13 has 13 groups both relate to occupation. From Figure 1.4 it is clear that occupation is a potentially influential variable on asthma rate.

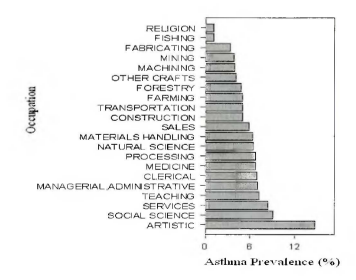


Figure 1.4 Asthma prevalence among occupations.

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The ARTISTIC, SOCIAL-SCIENCE, and SERVICES occupations show the highest asthma rates; FISHING and RELIGION have the lowest rate. Note, however, that some of these occupation groups are much smaller than others.

Physical exercise or physical activities can cause or worsen asthma symptoms (Ree and Kanabar, 2000, p. 16; Tan and Spector, 2002). A few people seem to get asthma attacks only when they run or take other exercise. This 'exercise-induced asthma' is especially a problem for young people. Exercise is just one of many things which show that the air passages are irritable in asthma. Cold air, tobacco smoke (e.g. in a pub), emotional stress, infections (such as colds and flu), sulphur dioxide (used as a preservative in soft drinks and wine) can all be triggers of asthma or asthma attacks (AAIR, 2003). The survey data also support this viewpoint. People who answered yes to the question of whether they participate in leisure time activity have an overall higher asthma rate.

For PAC6DFD which describe whether respondents participate in daily physical activities more than 15 minutes, people who answer yes have an asthma rate of 8.59% while people answer no have an asthma rate of 6.71%. The same pattern exists in other variables such as physical activity index.

For the demographic environment variables such as SDC6GCB (country of birth), SDC6GRAC (race or color), DHC6GMAR (marital status), the frequency tables generated by SAS show that respondents born in Canada have a higher asthma rate (8.6%) than people born in USA or Europe (5.4%), respondents born in Asia have an asthma rate as low as 1.7%, and white people have a higher asthma rate (8.01%) than other races (6.12%).

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Marital status is another demographic variable we are interested in. For the three category groups in this survey, we find that single persons have the highest asthma rate of 10.67%, followed by widow/separate/divorced person at 7.94% and married/living with a common law/partner at only 5.44%.

Smoking is one of the risk factors often thought to increase the risk of developing asthma. For individuals with asthma, smoking can make symptoms worse. In our study we find individuals who smoke daily have an asthma rate of 8.01%, higher than the overall rate of 7.81%. For those who smoke, variable SMC6_3 provides information about the starting age of their smoking habit. From Figure 1.5 we see that individuals with asthma start smoking relatively earlier than healthy ones.

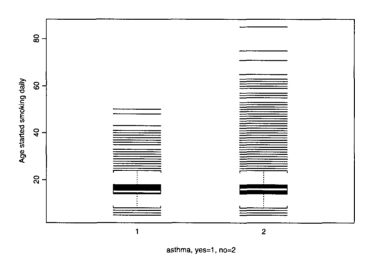


Figure 1.5 Boxplot of respondents' age of started smoking daily.

But for those who smoking daily, there seems to be little difference on the number of cigarettes smoked daily between the asthma group and non-asthma group (see Figure 1.6).

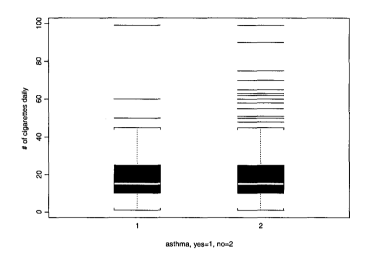
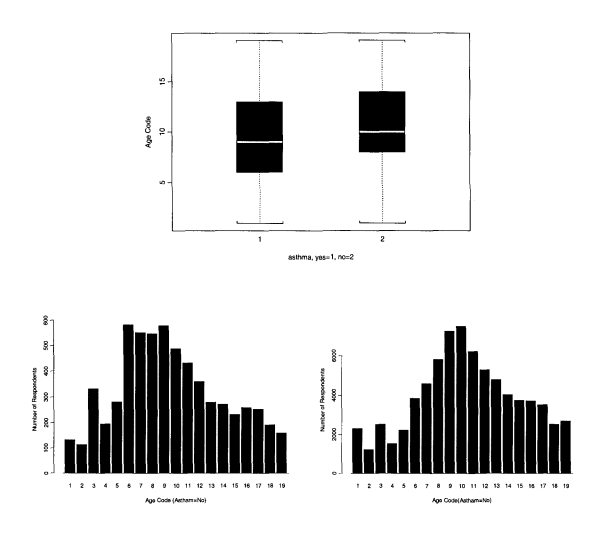


Figure 1.6 Boxplot of the number of cigarettes smoked per day.

Indoor air is another important factor that influences asthma prevalence. Research results from the Canadian Human Activities Pattern Survey (CHAPS) indicate that adults in Canada spend about 90% of their time indoors (Reports on Respiratory Disease in Canada of CHAPS, 2003, p. 27). As multiple concomitant exposures may heighten sensitivities, a combination effect is important to consider when indoor air exposures are suspected to be a cause of illness. Our data show that people who own the dwelling they live in have a lower asthma rate of 7.08%, by contrast the rest have asthma rate of 9.81%. The number of bedrooms also shows some effect on asthma rate. People living in dwellings with 1, 2, 3, 4 bedrooms have asthma rate of 8.91%, 8.41% 7.38% and 7.48%, respectively. People in households with 1, 2, 3, 4, 5 persons living in has asthma rate of 7.73%, 7.58%, 7.71%, 8.18% and 7.76%, respectively. Grouping people by whom they are living with reveals that single parents with children living together have the highest asthma rate of 12.68%.

The last issue we explore here is about the effect of basic characteristics of respondents on asthma prevalence.



Age Code	Age Range	Age Code	Age Range	Age Code	Age Range
1	0 TO 3 YEARS	8	25 TO 29 YEARS	15	60 TO 64 YEARS
2	4 TO 5 YEARS	9	30 TO 34 YEARS	16	65 TO 69 YEARS
3	6 TO 9 YEARS	10	35 TO 39 YEARS	17	70 TO 74 YEARS
4	10 TO 11 YEARS	11	40 TO 44 YEARS	18	75 TO 79 YEARS
5	12 TO 14 YEARS	12	45 TO 49 YEARS	19	80 YEARS OR OLDER
6	15 TO 19 YEARS	13	50 TO 54 YEARS		
7	20 TO 24 YEARS	14	55 TO 59 YEARS		

Figure 1.7 Age distribution of respondents who have asthma and who do not have asthma.

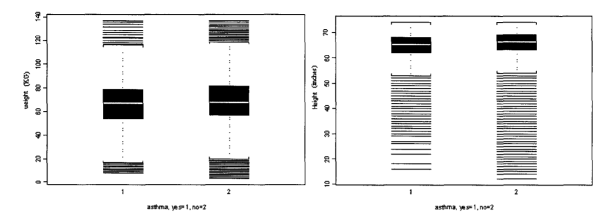


Figure 1.8 Boxplots of respondents' weight and height.

From Figure 1.7 we see that individuals with asthma have a relative young age. SAS results show that asthma prevalence is different by gender. From boxplots in Figure 1.8 we see that Weight and Height seem to have little effect on asthma prevalence. A look at the distribution by gender shows that females have a higher asthma prevalence than males, there respective prevalence rates are 8.47% and 7.16%. Our calculations also show that individuals with the airway diseases of chronic bronchitis or emphysema the asthma prevalence soar as high as 33.87%.

Chapter 2: Contingency Table Analysis for the Selected Variables in the NPHS

2.1 Contingency Table Analysis

Contingency tables are used to classify possible values of two or more variables. In each cross classified cell the number of observations is reported. They can be viewed as joint frequency distributions of variables.

We focus on the case of two variables: a row variable (R) and a column variable (C). Let R vary by a row index i, where i=1,2,...I, and let C vary by a column index j, where j=1,2,...J. It is customary to let R denote the explanatory variable and C the outcome variable. We denote the cell frequency of the ith row and the jth column by f_{ij} and the respective expected frequency by F_{ij} . See Table 2.1.

		Respondent variable				
		Level 1	Level 2	•••	Level J	Total
Explanatory Variable	Level 1	f ₁₁ (F ₁₁)	$f_{12}(F_{12})$		$f_{1J}\left(F_{1J}\right)$	$f_{1+}(F_{1+})$
	Level 2	$f_{21}(F_{21})$	f ₂₂ (F ₂₂)	•••	$f_{2J}\left(F_{2J}\right)$	$f_{2+}(F_{2+})$
	Level I	f ₁₁ (F ₁₂)	f ₁₂ (F ₁₂)		f _{IJ} (F _{IJ})	$f_{I+}\left(F_{I+}\right)$
	Total	f ₊₁ (F ₊₁)	f ₊₂ (F ₊₂)	•••	f _{+J} (F _{+J})	f++ (F++)

 Table 2.1
 Contingency table structure.

A widely used test statistic for testing the independence model is the Pearson's χ^2 statistic. Denote by \hat{F}_{ij} , the estimated expected frequency under independence,

 \hat{F}_{ij} = $f_{i+}\,f_{+j}$ / $f_{++}\,$, then Pearson's χ^2 statistic can be shown to be:

$$\chi^{2} = \sum_{i=1}^{I} \sum_{j=1}^{J} (\hat{F}_{ij} - f_{ij})^{2} / \hat{F}_{ij} . \qquad (2.1)$$

It can be proved (see Agresti, 1990, p. 44) that this statistic has an asymptotically chisquared distribution with degrees of freedom equal to (I-1)(J-1) when the hypothesis of independence is true.

In the contingency table analysis we also can use the likelihood ratio method. This method leads to another widely used statistic in contingency table analysis, the log likelihood test statistic G^2 given by:

$$G^{2} = -2\log Q = 2\sum_{i=1}^{I} \sum_{j=1}^{J} f_{ij} \log(f_{ij} / \hat{F}_{ij}) . \qquad (2.2)$$

 G^2 is asymptotically distributed as χ^2 under the independent hypothesis. The degrees of freedom can be calculated as the difference between the parameters under the unrestricted model (IJ-1) and the number of parameters under the restricted model (I-1)+(j-1) resulting in (I-1)(J-1) (Powers and Xie, 2000, pp. 88-105).

2.2 Application of Contingency Table Analysis to the NPHS Data

Using the programs listed in Appendix C we test for independence between the status of asthma (CCC6_1C) and each of the variables selected for study. The results are listed in Appendix B. Examining the P-values for both the Pearson method and the likelihood

ratio method (LLR), all the variables show a significant association with asthma except for: GE36GCMA (Derived 1991 Census Metropolitan Area - grouped), DHC6GHSZ (Derived household size - grouped) and DHC6GLE5 (Derived persons <= 5 years old in household - grouped). This suggests that most of the variables we selected from the source data set are potentially useful to explain asthma prevalence. That is, there is a significant difference in distribution between asthma and non-asthma groups for these variables.

Note that a total of 30 Chi-square statistical tests are reported in Appendix B. This means the overall probability of making the Type I error may be large. Note, however, that the P-values reported are very small for the most part (<0.0001) due to the large sample sizes. Thus, the single-test level of significance α could be made sufficiently small (much smaller than 5%) to offset the overall Type I error probability.

To see this in detail, consider m independent tests with a common significance level α . The overall significance level is:

 $P_m(\alpha)=P(rejecting independence in at least one test | all tables are independent)$. Clearly $P_m(\alpha)=1-(1-\alpha)^m$. Table 2.2 shows that, for m=30 tests, $P_m(\alpha)<0.05$ if $\alpha<0.0017$.

Table 2.2Values of the overall Type I error $P_m(\alpha)$

$P_m(\alpha)$
0.0030
0.0149
0.0498
0.1448

for individual significance levels α and m=30.

2.3 Test for Trend

Many of the variables contained in the data set are ordinal scaled. We are interested in whether there is a discernible trend in the level of association between the factor and the asthma prevalence rate. The trend test used in this section addresses this question and is very useful for two-way contingency tables with binomial response variables. The test is sensitive to the linearity between response variable and experimental variables and detects trends that would otherwise not be noticed by more crude methods. It is based upon the regression coefficient for the weighted linear regression of the binomial proportions on the scores of the levels of the explanatory variable. See Margolin (1988), Agresti (1990), and Stokes *et al.* (2000).

For I × 2 tables with ordered rows, using the linear probability model, for each row let $y_{1|I}$ denote the probability of response 1, and let $p_{1|i}$ denote the sample proportion, i = 1, ..., I. Let { x_i } be scores assigned to the rows. For the linear probability model

$$y_{1i} = \alpha + \beta x_i \quad . \tag{2.3}$$

the ordinary least squares fit gives the prediction equation

$$\hat{y}_{1i} = p_{+1} + b(x_i - \overline{x})$$
 (2.4)

where

$$\overline{\mathbf{x}} = \frac{\sum (n_i + x_i)}{n} , \quad \mathbf{b} = \frac{\sum [n_i + (p_{1|i} - p_{+1})(x_i - \overline{\mathbf{x}})]}{\sum [n_i + (x_i - \overline{\mathbf{x}})^2]} . \quad (2.5)$$

The Cochran-Armitage asymptotical standard normal statistic for trend is then given by

$$z = \left\{ \frac{b^2}{p_{+1}p_{+2}} \sum n_i + (x_i - \overline{x})^2 \right\}^{1/2} .$$
 (2.6)

When the linear probability model holds, the statistic z, tests for a linear trend in the proportions. The trend test may give strong evidence regarding positive or increasing linear trends, constant or stable trends over time, and negative or decreasing trends. SAS computes one-sided and two-sided p-values for the trend test (See SAS/STAT User's Guide Version 7 and Manitoba Centre for Health Policy and Evaluation, 1999). When the test statistic is greater than its expected value of zero, SAS computes the right-sided p-value, which is the probability of a larger value of the statistic occurring under the null hypothesis. A small right-sided p-value supports the alternative hypothesis of increasing trend in column 1 probability from row 1 to row I. When the test statistic is less than or equal to zero, SAS computes the left-sided p-value. A small left-sided p-value supports the alternative of decreasing trend (see SAS/STAT User's Guide Version 7).

Table 2.2 summarizes the z value obtained from the SAS analysis (programs is listed in the Appendix):

Variable	Cochran-Armitage Trend Test		
DHC6GAGE	Statistic (Z) -24.3017		
(Age – grouped)	One-sided Pr < Z <.0001		
	Two-sided Pr > Z <.0001		
DHC6GBED	Statistic (Z) 1.2591		
(Derived number of bedrooms in dwelling -	One-sided Pr > Z 0.1040		
grouped)	Two-sided $Pr > Z = 0.2080$		
PAC6DFR	Statistic (Z) -5.9025		
(Derived frequency of all physical activity)	One-sided Pr < Z <.0001		
	Two-sided Pr > Z <.0001		

Table 2.2Results of trend test.

Variable	Cochran-Armitage Trend Test			
SMC6_4	Statistic (Z) -3.8102			
(Number of cigarettes smoked each day - daily	One-sided Pr < Z <.0001			
smoker)	Two-sided $Pr > Z = 0.0001$			
SMC6_4(<=15)	Statistic (Z) -1.1955			
(Number of cigarettes smoked each day<=15	One-sided Pr < Z 0.1159			
cigarettes)	Two-sided $Pr > Z = 0.2319$			
SMC6_4(>15)	Statistic (Z) -2.0695			
((Number of cigarettes smoked each day>15	One-sided Pr < Z 0.0193			
cigarettes)	Two-sided Pr > $ Z $ 0.0385			
HWC6GHT	Statistic (Z) -15.3412			
(Height - adults and children – grouped)	One-sided Pr < Z <.0001			
	Two-sided $Pr > Z < .0001$			
HWC6G3KG	Statistic (Z) -15.0593			
(Derived standard weight – grouped-all	One-sided Pr < Z <.0001			
respondents)	Two-sided Pr > Z <.0001			
НЖС6G3КG	Statistic (Z) 2.3580			
(Derived standard weight – grouped-for respondents	One-sided Pr > Z 0.0092			
weights more than 70kg)	Two-sided $Pr > Z = 0.0184$			
HWC6GBMI	Statistic (Z) 6.0396			
(Derived Body Mass Index – grouped, respondents	One-sided Pr > Z <.0001			
aged 20~64 years old)	Two-sided $Pr > Z < .0001$			
SMC6 3	Statistic (Z) -7.2516			
(Age started smoking daily)	One-sided Pr < Z <.0001			
	Two-sided Pr > Z <.0001			

From Table 2.3 we see that at the α =5% level of significance there are decreasing trends in age, height, and age at smoking start to the prevalence of getting asthma. There is no trend between number of bedrooms, energy expenditure and asthma. Test to all respondents show that there is a decreasing trend between the number of cigarettes smoking daily to the prevalence of getting asthma. But if we look at the data we find that for those who smoke less than 15 cigarettes everyday there is no trend, trend only exists in respondents who smoke more than 15 cigarettes everyday. Body mass index shows

that for respondents who are 20 to 64 years old there is an increasing trend between the value of the index and asthma prevalence. Study of the variable of respondents' weight also shows this trend for the same age group. There is also increasing trend in derived standard weight.

In summary up the above results, we conclude that the younger the respondents the higher the asthma prevalence. Smoking has an adverse effect on respondents: the early they start smoking daily the larger the asthma prevalence. The decreasing trend between cigarettes smoked daily and asthma prevalence seems peculier at first. But we should consider the fact that the elder and longer a person smokes, the larger the potential the person smokes more cigarettes daily. For those who smoke more there may be some adaptation developed which helps them to defend against asthma. It is also true that people who suffer asthma tend to smoke less if they do have the smoking habit. Asthma tends to be suffered by those who are young and whose respiratory systems are not getting used to the smoking environment.

Chapter 3: Odds Ratio and Relative Risk Analysis

3.1 Odds Ratio and Relative Risk

It is helpful to describe the chances that a binary response variable leads to a success in terms of the odds of that event. The odds of success is defined to be the ratio of the probability of a success to the probability of a failure. Thus if p is the true success probability, the odds of success is p/(1-p).

	Yes	No	Total	Proportion of Yes
Group 1	n ₁₁	n ₁₂	n ₁₊	$\hat{p}_1 = n_{11}/n_{1+}$
Group 2	n ₂₁	n ₂₂	n ₂₊	$\hat{p}_2 = n_{21}/n_{2+}$
Total	n ₊₁	n ₊₂	n	

Table 3.1 A hypothetical 2×2 table.

For the 2×2 table displayed in Table 3.1, the odds ratio of interest is the one that compares the odds of the Yes proportion for Group 1 to the odds of the Yes proportion for Group 2. Specifically, denote the odds ratio as O_R , thus its estimate is

$$\hat{O}_{R} = \frac{\hat{p}_{1}/(1-\hat{p}_{1})}{\hat{p}_{2}/(1-\hat{p}_{2})} = \frac{n_{11}n_{22}}{n_{12}n_{21}}, \qquad (3.1)$$

where p_1 is the asthma probability in group 1 and p_2 is the asthma probability in group 2 whose estimates are \hat{p}_1 and \hat{p}_2 . The odds ratio ranges from 0 to infinity. When O_R is 1, there is no association between the row variable and the column variable. When O_R is greater than 1, group 1 is more likely than group 2 to have a Yes response and vise versa. It is convenient to take log of odds ratio, thus we get:

$$f = \log\{\hat{O}_{R}\} = \log\left\{\frac{\hat{p}_{1}/(1-\hat{p}_{1})}{\hat{p}_{2}/(1-\hat{p}_{2})}\right\} = \log\{\hat{p}_{1}/(1-\hat{p}_{1})\} - \log\{\hat{p}_{2}/(1-\hat{p}_{2})\} \quad .$$
(3.2)

The estimate of the variance of f is

$$\mathbf{v}_{f} = \left\{ \frac{1}{\mathbf{n}_{11}} + \frac{1}{\mathbf{n}_{12}} + \frac{1}{\mathbf{n}_{21}} + \frac{1}{\mathbf{n}_{22}} \right\}$$
(3.3)

so that an approximate 100(1- α)% confidence interval for O_R can be written as exp(f $\pm z_{\alpha/2}\sqrt{v_f}$).

Relative risk is a commonly used quantity in epidemiological studies. Relative risk is the risk of developing a particular condition (often a disease) for one group compared to another group. For data collected prospectively, the relative risk is written as $R_R=p_1/p_2$, thus from the definition of odds ratio and relative risk it is true that

$$\hat{R}_{R} = \hat{O}_{R} \times \frac{\{1 + (n_{21}/n_{22})\}}{\{1 + (n_{11}/n_{12})\}} \quad .$$
(3.4)

For cross sectional data, the quantity \hat{p}_1/\hat{p}_2 does not indicate risk since the disease and risk factor are assessed at the same time. When n_{11} and n_{21} are small relative to n_{12} and n_{22} , that is, the rare outcome assumption holds, \hat{O}_R is approximately equal to \hat{R}_R . Usually the outcome of interest needs to occur less than 10% of the time for \hat{O}_R and \hat{R}_R to be similar. For our data we have average asthma rate of 7.81% so it is proper for us to use \hat{O}_{R} as an approximation to \hat{R}_{R} in many cases. For details, see Stokes *et al.* (2000, pp. 29-33).

3.2 Application of Odds Ratio Analysis to the NPHS Data

To calculate odds ratio estimates and confidence intervals we just need to combine and adjust variables levels to get 2×2 tables. Thus there will be many odds ratios. The odds ratio results for those we are most interested in appear in Table 3.2. The last column contains a test for H₀: \hat{O}_R =1.

Test Meaning	Case-Control (Odds Ratio)	99% CI	χ^2 P-value
Respondents who attend physical activity daily VS not (PAC6DFD)	1.3062	1.2065 1.4141	Pearson 75.4897 <.0001 LLR 73.4328 <.0001
Respondents who attend physical activity ACTIVLLY VS not (PAC6DPAI)	1.3187	1.2074 1.4403	Pearson 65.6666 <.0001 LLR 62.7610 <.0001
Respondents who were born in America & Europe VS other country (SDC6GCB)	0.6511	0.5687 0.7454	Pearson 67.7489 <.0001 LLR 74.5920 <.0001
Respondents who were born in Asia VS other country (SDC6GCB)	0.1957	0.1429 0.2682	Pearson 220.6924 <.0001 LLR 311.3942 <.0001
Respondents who were born in Canada VS other country (SDC6GCB)	2.2713	2.0200 2.5539	Pearson 341.9383 <.0001 LLR 397.9497 <.0001
Respondents who are white VS all others (SDC6GRAC)	1.3356	1.1847 1.5058	Pearson 626.7386 <.0001 LLR 613.3712 <.0001
Respondents who are single VS all others (SDC6GMAR)	1.9125	1.7874 2.0463	Pearson 626.7386 <.0001 LLR 613.3712 <.0001
Respondents who attend physical activity regular VS not (PAC6DFR)	1.1906	1.1005 1.2880	Pearson 32.6571 <.0001 LLR 32.9934 <.0001
The province with the highest asthma rate vs the lowest one (Nova Scotia VS Newfoundland, PRC6_CUR)	1.5062	1.0785 2.1035	Pearson 10.0911 0.0015 LLR 10.4245 0.0012

Table 3.2 Results of odds ratio analysis.

Test Meaning	Case-Control (Odds Ratio)	99% CI	χ^2 P-value
The province with the asthma rate high than average vs the lower ones (Nova Scotia & Ontario & Quebec VS all others ,PRC6_CUR)	1.0975	1.0217 1.1789	Pearson 11.2069 0.0008 LLR 11.3012 0.0008
The rural AREA vs the urban area (GE36GURB)	0.8166	0.7129 0.9354	Pearson 14.8099 0.0001 LLR 15.1325 0.0001
The Occupation with the highest asthma rate vs the lowest one: (Personal service VS Agriculture, LFC6GI13)	2.5639	1.6839 3.9039	Pearson 35.5754 <.0001 LLR 41.6143 <.0001
The Occupation with the higher than average asthma rate vs the lower ones: (Finance, Community Services & Personal service VS all other occupations, LFC6G113)	1.5160	1.3692 1.6786	Pearson 111.9838 <.0001 LLR 108.2655 <.0001
Children(age <12) VS Adult (PAC6DEE)	1.6247	1.4929 1.7683	Pearson 221.8114 <.0001 LLR 201.6865 <.0001
Respondents who attend leisure time physical activity VS not (PAC6DLEI)	1.1474	1.0088 1.3051	Pearson 7.5770 0.0059 LLR 7.8128 0.0052
Respondents who are widowed, separated, divorced VS married/common-law/partner (SDC6GMAR)	1.5006	1.3342 1.6878	Pearson 80.314 <.0001 LLR 74.1895 <.0001
Respondents who smoke vs not (SMC6_3)	1.0338	0.9512 1.1236	Pearson 1.0579 0.3037 LLR 1.0520 0.3050
Respondents who start smoke daily before 20 years old vs start after 20 years old (SMC6_3)	1.4853	1.1949 1.8463	Pearson 22.1964 <.0001 LLR 23.8456 <.0001
Respondents who smoke daily more than 10 cig. vs less than 10 (SMC6_4)	0.7644	0.6515 0.8968	Pearson 18.8569 <.0001 LLR 18.8569 <.0001
Respondents who smoke daily more than 10 cig & less than20 years old. vs less than 10 & less than20 years old (SMC6_4)	1.2247	0.8577 1.7486	Pearson 2.1530 0.1423 LLR 2.1461 0.1429
Respondents who smoke daily more than 10 cig & elder than 20 years old. vs less than 10 & elder than 20 years old (SMC6_4)	0.8270	0.6876 0.9946	Pearson 7.0467 0.0079 LLR 6.8573 0.0088
Dwelling not owned by respondents' family member vs yes (DHC6_OWN)	1.4263	1.3276 1.5323	Pearson 164.2046 <.0001 LLR 157.2275 <.0001
Dwelling have no more than 2 bedrooms vs yes (DHC6GBED)	1.1458	1.0633 1.2347	Pearson 22.0446 <.0001 LLR 21.6534 <.0001
FEMALE vs MALE (DHC6_SEX)	1.2036	1.1251 1.2875	Pearson 50.2127 <.0001 LLR 50.2966 <.0001

Test Meaning	Case-Control (Odds Ratio)	99% CI		χ^2 P-value
FOR CHILDREN(<12 YEARS OLD) MALE vs FEMALE (dhc6_sex)	1.6211	1.3882 1.8931	Pearson LLR	65.2512 <.0001 65.8816 <.0001
FOR ADULT (AGE>=12 YEARS OLD) FEMALE vs MALE (DHC6_SE)	1.4311	1.3261 1.5445	Pearson LLR	147.9869 <.0001 148.8596 <.0001
Adult Respondents who are overweight vs not (HWC6GSW)	1.1524	1.0399 1.2771	Pearson LLR	12.6688 0.0004 12.4570 0.0004
Respondents who have chronic bronchitis or emphysema vs not (CCC6_1H)	6.7796	6.0304 7.6218	Pearson LLR	2316.2080 <.0001 1391.1395 <.0001

The results in Table 3.2 indicate that the odds ratios significantly depart from 1 in most cases considered. The results go along with the features noted in the exploratory analysis in Section 1.4. For instance, a significantly higher asthma prevalence occurs for respondents born in Canada compared to those born in other countries. A significantly higher asthma prevalence occurs among those who are single compared to those in other marital circumstances. Also a significantly higher asthma prevalence occurs among females when compared to males. On the opposite side, living in rural areas significantly decreases the asthma prevalence when compared to urban areas. Respondents living in dwellings owned by their family members have relatively lower asthma prevalence.

Chapter 4 Loglinear and Logistic Regression Analysis

4.1 Loglinear Regression Models for Two-way s×r Tables and High Dimension Tables

When a sample of n observations is classified with respect to two categorical variables, one having s levels and the other having r levels, then the resulting frequencies can be displayed in an s×r contingency table, as shown below. The corresponding cell probabilities are π_{ij} , with row and column marginal probability π_{i+} and π_{+j} , respectively.

Level of X		Total			
	1	2	••••	r	Total
1	n ₁₁	n ₁₂	•••	n _{1r}	n ₁₊
2	n ₂₁	n ₂₂		n _{2r}	n ₂₊
		•••	•••		
S	n _{s1}	n _{s2}		n _{sr}	n _{s+}
Total	n ₊₁	n ₊₂	•••	n _{sr}	n

Table 4.1Representation of an s×r table.

The generalization of the saturated loglinear model for this s×r contingency table is:

$$\log(m_{ij}) = \mu + \lambda_i^{X} + \lambda_j^{Y} + \lambda_{ij}^{XY}, \qquad i = 1,...,s; j = 1,...,r , \qquad (4.1)$$

where $m_{ij}=n \pi_{ij}$ is the expected frequency in the (i,j)th cell. The parameter μ is fixed by the sample size n and the model has s+r+sr parameters. The parameters in the above model satisfy the sum-to zero constraints:

$$\sum_{i=1}^{s} \lambda_{i}^{X} = 0, \qquad \sum_{j=1}^{r} \lambda_{j}^{Y} = 0, \qquad \sum_{i=1}^{s} \lambda_{ij}^{XY} = \sum_{j=1}^{r} \lambda_{ij}^{XY} = 0.$$
(4.2)

The constraints implies that there only (s-1)+(r-1)+(s-1)(r-1)=sr-1 parameters can be estimated and zero df for testing lack of fit. Letting $\hat{m}_{ij} = n_{i+}n_{+j}/n$, the likelihood ratio statistic

$$G^{2} = 2\sum_{i=1}^{s} \sum_{j=1}^{r} n_{ij} \log(n_{ij} / \hat{m}_{ij}) . \qquad (4.3)$$

test the hypothesis $H_0: \lambda_{ij}^{XY} = 0$, for i=1,...s-1, j=1,...r-1. Under the null hypothesis of independence, G^2 has an approximate Chi-square distribution with (s-1)(r-1) df. If H_0 is true, the reduced model $\log(m_{ij}) = \mu + \lambda_i^X + \lambda_j^Y$ is the model of independence of X and Y. This model has (s-1)+(r-1) linearly independent parameters of λ and (s-1)(r-1) df for testing lack of fit.

The same logic holds for loglinear models dealing with high dimension tables. For a W way contingency table we can consider all the combination of W-1, ...2 factor interaction terms and all main effects. As the number of dimensions of a contingency table increases, there will be some complicating factors. The number of possible interaction parameters will increase tremendous. This will result in hundreds of or even thousands of parameters to be estimated. Even if the sample size is large, whether these huge number of parameters can be estimated is a problem, since there may be many observed cell counts equal to zero. There may even be marginal totals equal to zero. For details see Stokes *et al.* (2000, pp. 560-568).

In our data set there are 32 variables, some of them having as many as 66 levels. So to construct a loglinear model we need to include only those important variables and combine variable levels to get a reasonable model. Thus we have very a large number of models to chose. All our variables as described in Chapter 1 can be divided into 7 groups. Variables from the same group describe similar or related aspects of respondents. So we chose about one variable from each group. These selected variables all have significant Pearson chi-square and log likelihood ratio chi-square values. Another reason for selecting only one variable from each group is to avoid the multicolinearity problem that arises when highly correlated variables are included. These variables are: CCC6_1C (Has asthma); PRC6_CUR (Province of residence); LFC6GI13 (Industry Codes for main job -13 groups); PAC6DPAI (Derived physical activity index); SMC6_3 (Age started smoking daily - daily smoker); DHC6_OWN (Dwelling owned by household member); DHC6_SEX (Gender). If we consider all the 6-factor interactions and all its lower order interaction as well as main effects, then we will have thousands of possible parameters to estimate. That is a huge but unnecessary task. We deal with this problem with two methods.

One is to combine the levels of the variables. We combine the 10 levels of province to two: the first level indicating higher asthma prevalence rate, which include Ontario Quebec and PEI. The second level contains all other provinces. We also do a grouping of the 13 levels of occupation. We put those with no occupation in the first level, those occupations with high asthma rate (Finance, community service, personal services, transportation, retail trade, business services, public administration) in the second level and all other occupation in the last level.

The other alternative is to consider hierarchical models. Our aim in constructing loglinear models is to capture the relations among variables. We focus on hierarchy models since non hierarchy models have little meaningful interpretation. By fitting hierarchical models we find the best model which only includes significant interaction terms. We apply the following strategy: we first fit models with all 6-factor interactions, if it is not significant we will try the 5-factor interactions models and keep on till we could confirm that the lower than the present order model which are significant. Then we delete all the terms which are not significant in the present model to get the best one.

4.2 Results of the Seven-way Loglinear Regression Model

The results of the model fitting are listed in the Table 4.2. The 2-factor hierarchy model indicates that not all variables have an effect on CCC6_1C. But the G² value is significant, which indicates that we shall consider adding more interaction terms to improve the model fitting. The 5-factor hierarchy model is significant, but the 6-factor one is not. The G^2 value increases from 0.0002 (df=1) for the 6-factor hierarchy model to 65.54 (df =14) for the 5-factor ones. After deleting all the non significant terms in the 6factor hierarchy model we get the best model which describes the relationship of all the variables. The terms included in the best model and their Chi-square values are listed in table 4.2. Comparing the 5 and 6 factor hierarchy models we see that for the six factor interaction terms only CCC*PRC*LFC*DHC*SMC*DHC6 & CC*LFC*PAC*DHC*SMC*DHC6 are significant and need to be added to the 5-factor hierarchy model to improve the fitting. Many of the 2 and 3 way interactions are significant in the 6 factor hierarchy model. For the three way interaction terms which are related to CCC6_1C, we see that CCC6_1C*PRC6_CU*LFC6GI13, CCC6_1C*PRC6_CU*PAC6DPAI, CCC6_1C*LFC6GI1*PAC6DPAI, CCC6_1C*LFC6GI1*DHC6_SEX, CCC6_*PRC6_*LFC6G*DHC6_S, CCC6_1C*PAC6DPA*DHC6_SEX, CCC6_1C*LFC6GI1*DHC6_OWN, CCC6_1C*PAC6DPA*DHC6_OWN are significant at 5% level.

Table 4.2 Results of loglinear model.

			Pr >
Source	DF	Chi-Square	ChiSq
CCC6_1C	1	3012.09	<.0001
PRC6_CUR	1	203.27	<.0001
LFC6GI13	2	877.27	<.0001
CCC6_1C*LFC6GI13	2	71.04	<.0001
PRC6_CUR*LFC6GI13	2	31.33	<.0001
CCC6_1C*PRC6_CU*LFC6GI13	2	12.37	0.0021
PAC6DPAI	2	1053.14	<.0001
CCC6_1C*PRC6_CU*PAC6DPAI	2	8.06	0.0178
CCC6_1C*LFC6GI1*PAC6DPAI	2	8.99	0.0112
DHC6_SEX	1	103.7	<.0001
CCC6_1C*DHC6_SEX	1	7.25	0.0071
LFC6GI13*DHC6_SEX	2	406.52	<.0001
CCC6_1C*LFC6GI1*DHC6_SEX	2	21.2	<.0001
PRC6_CU*LFC6GI1*DHC6_SEX	2	8.52	0.0141
CCC6_*PRC6_*LFC6G*DHC6_S	2	6.62	0.0366
PAC6DPAI*DHC6_SEX	2	210.34	
CCC6_1C*PAC6DPA*DHC6_SEX	2	6.4	0.0408
PRC6_CU*PAC6DPA*DHC6_SEX	2	1.41	0.494
PRC6_*LFC6G*PAC6D*DHC6_S	2	22.11	<.0001
CCC6*PRC6*LFC6*PAC6*DHC6	2	22.95	<.0001
SMC6_3	1	1016.43	<.0001
PRC6_CUR*SMC6_3	1	4.27	0.0388
LFC6GI13*SMC6_3	2	59.17	<.0001
PRC6_CUR*LFC6GI13*SMC6_3	2	9.55	0.0084
CCC6_*PRC6_*LFC6G*SMC6_3	2	10.97	0.0042
PAC6DPAI*SMC6_3	1	7.64	0.0057
CCC6*PRC6*LFC6*PAC6*SMC6	2	7.55	0.023
CCC6_1C*DHC6_SEX*SMC6_3	1	25.26	<.0001
PRC6_CUR*DHC6_SEX*SMC6_3	1	18.83	<.0001
CCC6_*PRC6_*DHC6_*SMC6_3	1	24.96	
LFC6GI13*DHC6_SEX*SMC6_3	2	18.99	<.0001
CCC6_*LFC6G*DHC6_*SMC6_3	2		<.0001
CCC6*PRC6*LFC6*DHC6*SMC6	2	10.2	0.0061
CCC6_*PAC6D*DHC6_*SMC6_3	1	4.46	0.0347
LFC6G*PAC6D*DHC6_*SMC6_3	2	8.46	0.0145
CCC6*LFC6*PAC6*DHC6*SMC6	2	7.44	0.0242
DHC6_OWN	1	435.56	<.0001
PRC6_CUR*DHC6_OWN	1	4.34	0.0371
LFC6GI13*DHC6_OWN	2	18.41	0.0001
CCC6_1C*LFC6GI1*DHC6_OWN	2	6.48	0.0392
PRC6_CU*LFC6GI1*DHC6_OWN	2	9.33	0.0094
CCC6_*PRC6_*LFC6G*DHC6_O	2	2.04	0.36
PAC6DPAI*DHC6_OWN	2	30.68	<.0001
CCC6_1C*PAC6DPA*DHC6_OWN	2	12.28	0.0022
DHC6_SEX*DHC6_OWN	1	4.09	0.043

LFC6GI1*DHC6_SE*DHC6_OWN	2	6.4	0.0408
CCC6_*LFC6G*DHC6_*DHC6_O	2	7.12	0.0284
PRC6_*LFC6G*DHC6_*DHC6_0	2	6.54	0.038
LFC6G*PAC6D*DHC6_*DHC6_0	2	18.19	0.0001
CCC6*LFC6*PAC6*DHC6*DHC6	2	19.12	<.0001
SMC6_3*DHC6_OWN	1	152.52	<.0001
CCC6_*LFC6G*SMC6_*DHC6_0	2	9.88	0.0072
PRC6_*LFC6G*SMC6_*DHC6_0	2	0.17	0.92
PAC6DPAI*SMC6_3*DHC6_OWN	1	11.81	0.0006
CCC6_*PAC6D*SMC6_*DHC6_0	1	4.66	0.0309
PRC6_*PAC6D*SMC6_*DHC6_0	1	6.18	0.0129
LFC6G*PAC6D*SMC6_*DHC6_0	2	9.62	0.0081
DHC6_SEX*SMC6_3*DHC6_OWN	1	6.02	0.0141
CCC6_*DHC6_*SMC6_*DHC6_O	1	4.57	0.0326
PRC6_*DHC6_*SMC6_*DHC6_0	1	4.37	0.0365
CCC6*PRC6*DHC6*SMC6*DHC6	1	2.63	0.105
LFC6G*DHC6_*SMC6_*DHC6_0	2	14.1	0.0009
CCC6*LFC6*DHC6*SMC6*DHC6	2	16.81	0.0002
PRC6*LFC6*DHC6*SMC6*DHC6	2	12.41	0.002
CCC*PRC*LFC*DHC*SMC*DHC6	2	9.76	0.0076
LFC6*PAC6*DHC6*SMC6*DHC6	2	29.22	<.0001
CCC*LFC*PAC*DHC*SMC*DHC6	2	41.26	<.0001

4.3 Logistic, Probit & loglog Regression Models

Suppose we have N binary responses as independent Bernoulli radom variables. Let $x_i=(x_{i0}, x_{i1},..., x_{ik})$ denote the ith setting of values of k explanatory variables, i=1,2,...I, where $x_{i0}=1$. when more than one observation on Y occurs at a fixed x_i value, it is sufficient to record the number of observations n_i and the number of "1" outcomes. Thus let Y_i refer to this "success" count rather than to individual binary responses. The { Y_i i=1,...,I} are independent binomial random variables with $E(Y_i)=n_i\pi(x_i)$, where $N=n_1+...+n_I$. The joint probability mass function of $(Y_1,...,Y_I)$ is proportional to the product of I binomial terms, so we have:

$$\prod_{i=1}^{I} \pi(x_{i})^{y_{i}} [1 - \pi(x_{i})]^{n_{i} - y_{i}} = \left\{ \prod_{i=1}^{I} [1 - \pi(x_{i})]^{n_{i}} \right\} \left\{ \prod_{i=1}^{I} \exp\left[\log\left(\frac{\pi(x_{i})}{1 - \pi(x_{i})}\right)^{y_{i}}\right] \right\}$$
$$= \left\{ \prod_{i=1}^{I} [1 - \pi(x_{i})]^{n_{i}} \right\} \exp\left[\sum y_{i} \log\left(\frac{\pi(x_{i})}{1 - \pi(x_{i})}\right)\right] \quad .$$
(4.4)

Let $\pi(x) = \frac{\exp(\alpha + \beta x)}{1 + \exp(\alpha + \beta x)}$, after transformation we get $\log\left(\frac{\pi(x_i)}{1 - \pi(x_i)}\right) = \eta = \sum_j \beta_j x_{ij}$,

after some rearrangement we can write the log likelihood function as:

$$\mathbf{L}(\boldsymbol{\beta}) = \sum_{j} \left(\sum_{i} \mathbf{y}_{i} \mathbf{x}_{ij} \right) \boldsymbol{\beta}_{j} - \sum_{i} n_{i} \log \left[1 + \exp\left(\sum_{j} \boldsymbol{\beta}_{j} \mathbf{x}_{ij}\right) \right] \quad .$$
(4.5)

this depends on the binomial counts only through the sufficient statistics $\sum_{i} y_i x_{ij}$, j=0,...,k. By using a numeric procedure such as Newton-Raphson method we can get the ML estimate of β_i , thus we can get an estimate of $\pi(x_i)$.

Alternatively define $\pi(x) = \Phi[(x - \mu)/\sigma]$, where Φ is the standard normal cdf. If we use $\Phi^{-1}[\pi(x)] = \alpha + \beta x$ the model is the probit model. If we define $\pi(x) = \exp[-\exp(\alpha + \beta x)]$ and use $\log\{-\log[\pi(x)]\} = \alpha + \beta x$, we are fitting a complementary log-log model. All these three types of models are considered in our analysis. For details see Agresti (1990, pp. 102-115).

Our data set contains various kinds of variables. Categorical and continuous, ordinal and non ordinal all exist. There are also many structural missing values, such as variable SMC_3, which only indicates the starting age of respondents who smoke. Thus before we can fit the logistic regression model we need to rearrange the data.

For the categorical and non ordinal variables, the code values just indicate difference categories. It is not proper to put them directly to the design matrix of x. We need to create dummy variables for them. To fit our logistic and other two kinds of regression models, we use the reference code. That is, we select a base category, others effects are all compared to this base one. Parameter estimates using the reference coding scheme estimate the difference in the effect of each non-reference level compared to the effect of the reference level.

For ordinal variables we first look whether the increase of code value respondent to uniform change of the object. For example, the grouped variable DHC6GAGE. The variable describe the respondent's age. But we only get the grouped value due to confidentiality requirements. The one unit increase of code corresponds to an increase of 5 years. Thus for this variable we include it in the models using its own code value. Fortunately all the categorical ordinal variables are like this. Continuous variables such as PAC6DFM (monthly freq. of physical activity) are all included in the models directly.

It is a little complicated to treat the structural missing values. For example, variable LFC6GI13 describes the respondent's occupation. For those who do not work there will be at this variable if we include LFC6GI13 in the model. SAS will automatically delete cases with missing values when running a logistic regression procedure. Since we have many variables with structural missing values, if we do not rearrange the data then only a small number of cases can be left for us to do analysis. So for this variable we treat the respondents who do not have occupation as another category and make a dummy variable for it except the 12 dummy variables respondent to its levels. SMC6_3 (age start smoking) is another categorical variable with structural missing value due to respondents who do not smoke. If viewing those persons who do not smoke as they start smoking very late, say, in their 100 years old, then we could eliminate the structural missing value by substituting them with a large number say, 100. All the details about the rearranged codes and information about the design matrix are listed in Appendix A.

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4.4 Results of the Logistic, Probit, and Complementary Log-log Regression Models for All Respondents

The results in Tables 4.3-4.5 were obtained by using stepwise regression method in SAS. In addition to the log likelihood ratio test, parameters estimate, odds ratio estimates and some other statistics, we also do the Hosmer and Lemeshow goodness-of-fit test. Receiving operating characteristic curve (ROC) is also plotted using Splus.

Table 4.3Result of logistic regression.

Step 37. Effect DHC6DL5 entered:						
Model Convergence Status						
Convergence criterion (GCONV=1E-8) satisfied.						
	001		0		o, 500151100	-
			Model	Fit Statistic	cs	
			110401		Intercept	
				Intercept	and	
		Cri	terion	Only	Covariates	
		AIC		39793.480	38201.469	
		SC		39802.626	38512.448	
		-2	Log L	39791.480	38133.469	
			5			
		Tes	ting Global	Null Hypothes	sis: BETA=0	
	Test			hi-Square		> ChiSq
	Likelih	nood		1658.0108	33	<.0001
	Score			1547.9753	33	<.0001
	Wald			1440.1822	33	<.0001
			Residua	1 Chi-Square '	Fest	
		С	hi-Square	-	r > ChiSq	
			109.7621	32	<.0001	
NOTE: No (addit	ional) effec	cts n	net the 0.05	significance	level for en	try into the model.
		Anal	ysis of Max	imum Likeliho	od Estimates	
				Standard		
Р	arameter	\mathbf{DF}	Estimate	Error	Chi-Square	Pr > ChiSq
I	ntercept	1	-3.8002	0.1798	446.8662	<.0001
N	FLD	1	-0.3610	0.1140	10.0250	0.0015
N	В	1	-0.2023	0.0937	4.6595	0.0309
A	В	1	-0.1492	0.0500	8.8937	0.0029
G	E36GU1	1	-0.1730	0.0503	11.8512	0.0006
Н	WC6G1	1	-0.2285	0.0701	10.6107	0.0011
Н	WC6G2	1	-0.3293	0.0404	66.4116	<.0001
Н	WC6G3	1	-0.1705	0.0527	10.4460	0.0012
L	FC1	1	0.4606	0.0493	87.3481	<.0001
L	FC3	1	0.2319	0.0881	6.9243	0.0085
L	FC6	1	0.2472	0.0854	8.3843	0.0038
L	FC9	1	0.6246	0.0877	50.6819	<.0001
L	FC10	1	0.4770	0.0600	63.2354	<.0001
L	FC11	1	0.3843	0.0682	31.7653	<.0001
	FC12	1	0.2770	0.0740	14.0131	0.0002
	FC13	1	0.1982	0.0938	4.4629	0.0346
	AC6DPA1	1	0.5385	0.0586	84.5208	<.0001
	AC6DPA2	1	0.5149	0.0618	69.5356	<.0001
	AC6DPA3	1	0.4139	0.0602	47.2899	<.0001
	DC6G1	1	1.0289	0.1382	55.4054	<.0001
	DC6G2	1	0.8823	0.1477	35.7008	<.0001
	DC6G3	1	-0.7150	0.1943	13.5479	0.0002
S	MC6_31	1	-0.00140	0.000442	10.0113	0.0016

DHC6_02 DHC6_S2 DHC6GM2 DHC6GM3 DHC6GE1 DHC6GE4 DHC6GE5 DHC6GE6 DHC6GL5 DHC6DL5 DHC6DL6 DHC6GAGE	1 1 1 1 1 1 1 1 1 1	0.3001 0.1129 0.3870 0.5043 0.1448 0.3022 0.3616 0.3671 0.1592 0.3509 -0.0718	0.0333 0.0299 0.0535 0.0665 0.0499 0.0808 0.0692 0.0614 0.0784 0.0844 0.00662	14. 52. 57. 8. 13. 27. 35. 4. 17.	2565 2879 3058 5186 4107 9754 3423 7731 1234 2851 5309	<.0001 0.0002 <.0001 0.0037 0.0002 <.0001 <.0001 0.0423 <.0001 <.0001
		Odds Ra	atio Estima	ates		
	Effect NFLD NB AB GE36GU1 HWC6G1	Poin Estimat 0.69 0.80 0.84 0.84	te Cor 97 0 17 0 51 0 41 0 96 0	95% Wald nfidence I .557 .680 .781 .762 .694	Limits 0.871 0.982 0.950 0.928 0.913	
	HWC6G2 HWC6G3 LFC1	0.7: 0.84 1.58	43 0	.665 .760 .439	0.779 0.935 1.746	
	LFC3 LFC6	1.20	51 1 80 1	.061 .083	1.499 1.514	
	LFC9 LFC10 LFC11	1.8 1.6 1.4	11 1	.572 .433 .285	2.218 1.812 1.679	
	LFC12 LFC13	1.3	19 1	.141 .014	1.525	
	PAC6DPA1 PAC6DPA2	1.7:	74 1	.528	1.922	
	PAC6DPA3 SDC6G1	1.5		.344 .134	1.702 3.669	
	SDC6G2	2.43		.809	3.228	
	SDC6G3	0.4		.334	0.716	
	DHC6_02 DHC6_S2	1.3		.265	1.441	
	DHC6_52 DHC6GM2	1.1: 1.4		.056 .326	1.187 1.635	
	DHC6GM3	1.6		.453	1.886	
	DHC6GE1	1.1		.048	1.275	
	DHC6GE4	1.3	53 1	.155	1.585	
	DHC6GE5	1.43		.254	1.644	
	DHC6GE6	1.44		.280	1.628	
	DHC6DL5 DHC6DL6	1.1		.006	1.367	
	DHCODLO	1.42	20 1	.204	1.676	
Associa	tion of Pr	edicted Pr	robabilitie	es and Obs	served Res	ponses
	cent Disco		60.5 37.8	Gamma	0.2	
	cent Tied		1.6	Tau-a	0.0	
Pai	rs		337650512	с	0.6	
	Partitic		Hosmer and C6_1C = YES		v Test CCC6_10	- NO
Group	Total	Observe	_		served	Expected
1	6946		- ·	3.78	6695	6757.22
2	6934	35	50 296	5.75	6584	6637.25
3	6861			7.82	6486	6513.18
4	6907			4.89	6491	6512.11
5	6939			3.73	6496	6495.27
6 7	6916			0.72	6458	6415.28
8	6932 6932			9.97 3.64	6307 6217	6352.03
9	6962			0.45	6222	6248.36 6141.55
10	6989			2.47	6090	5906.53
			now Goodnes			
		Square	DF	ss-or-fit Pr > Chis		
		9.5752	8	<.000	-	

Table 4.4 Result for probit regression.

Model Convergence Status Convergence criterion (GCONV=1E-8) satisfied.

Model Fit Statistics

		Intercept
	Intercept	and
Criterion	Only	Covariates
AIC	39793.480	38213.265
SC	39802.626	38515.098
-2 Log L	39791.480	38147.265

Testing Gl	obal Null Hypothe	esis: BET	A=0
Test	Chi-Square	DF	Pr > ChiSq
Likelihood Ratio	1644.2145	32	<.0001
Score	1542.7256	32	<.0001
Wald	1460.9200	32	<.0001

Residual Chi-Square TestChi-SquareDFPr > ChiSq102.237431<.0001</td>

NOTE: No (additional) effects met the 0.05 significance level for entry into the model.

Association	of Predicted	Probabilities	and Observed	Responses
Percent	Concordant	60.5	Somers' D	0.227
Percent	Discordant	37.9	Gamma	0.230
Percent	Tied	1.6	Tau-a	0.032
Pairs		337650512	с	0.613

	Partition	for the Hos	mer and Lemesh	low Test	
		CCC6_1	C = YES	CCC6_1C	= NO
Group	Total	Observed	Expected	Observed	Expected
1	6964	246	183.41	6718	6780.59
2	6918	353	293.42	6565	6624.58
3	6917	377	352.17	6540	6564.83
4	6956	413	403.66	6543	6552.34
5	6906	454	450.18	6452	6455.82
6	6939	458	513.60	6481	6425.40
7	6926	623	592.18	6303	6333.82
8	6925	713	692.05	6212	6232.95
9	6931	755	819.28	6176	6111.72
10	6936	880	1053.74	6056	5882.26
	Hosmer an	nd Lemeshow	Goodness-of-Fi	t Test	

HODINCE GILG	Tierue Dirto M	00000000	,0 O T	110 1000
Chi-Squa	are	DF	Pr >	ChiSq
85.13	68	8		<.0001

Table 4.5 Result for Complementary log-log.

Model Convergence Status Convergence criterion (GCONV=1E-8) satisfied.

Mode	l Fit Statist	ics Intero	cept
	Intercept	and	f
Criterion	Only	Covaria	ates
AIC	39793.480	38199	.962
SC	39802.626	38510	.942
-2 Log L	39791.480	38131	.962
Testing Globa	1 Null Hypoth	esis: BETA	A=0
Test	Chi-Square	DF	Pr > ChiSq
Likelihood Ratio	1659.5177	33	<.0001
Score	1547.9753	33	<.0001
Wald	1459.6394	33	<.0001

Residual	Chi-Square	Т	est	t
Chi-Square	DF	Pr	>	ChiSq
102.8374	30		•	<.0001

NOTE: No (additional) effects met the 0.05 significance level for entry into the model.

Pairs

Association of Predicted Probabilitiesand Observed ResponsesPercent Concordant60.5Somers' D0.227Percent Discordant37.8Gamma0.231Percent Tied1.6Tau-a0.032

337650512

с

0.613

	Partition	for the Hosm	er and Leme	show Test	
		CCC6_1C	= YES	CCC6_10	2 = NO
Group	Total	Observed	Expected	Observed	Expected
1	6918	250	189.54	6668	6728.46
2	6901	348	296.24	6553	6604.76
3	6870	378	348.22	6492	6521.78
4	6902	420	393.83	6482	6508.17
5	6942	436	442.41	6506	6499.59
6	6917	455	498.63	6462	6418.37
7	6939	628	577.76	6311	6361.24
8	6922	715	679.45	6207	6242.55
9	6945	736	814.87	6209	6130.13
10	7062	906	1096.52	6156	5965.48

Hosmer and	Lemeshow	Goodnes	s-of-Fit Test
Chi-Squa	are	DF	Pr > ChiSq
92.6	327	8	<.0001

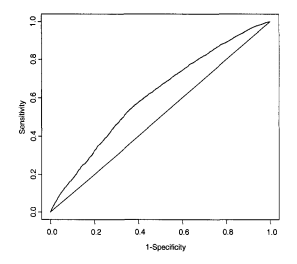


Figure 4.1 Receiving operating characteristic curve (ROC).

From the above results (refer to the programs and the variable meaning listed in the appendix A and H) we see that for all three types of regression the results are very similar. The likelihood ratio chi-square statistic are all large enough to justify the goodness of the model. But for all three models the Hosmer and Lemeshow goodness-of-fit Test shows some lack of fit in the models. Generally say, the variables included in the models are important and statistically significant, but there may still exist some other important factors to be included in the models. Consider the fact that our data set only contains the environmental factors thought to effect asthma prevalence but there could exist other important variables such as the genetic ones. It may be those variables that cause Hosmer and Lemeshow value in these models significant. In the next section we will apply an approach to alleviate this problem considering in fitting separate regression for infants, children and adults. We will also address problem of model validation.

To assess the ability of the logistic model to discriminate between events and nonevents the receiver operating characteristic (ROC) curve is a good diagnostic tool. For every respondent in the data set we can get the corresponding estimated probability of event. Sort these estimated probabilities and view every one of them as cut-points for predicting the response, then label any observation with an estimated event probability that exceeds or equals a given threshold (or cut-points) as being predicted to be an event. Calculate the proportion of event observations that were correctly predicted to have an event response, this is called the sensitivity corresponding to the cut-points. We can also get the relevant one minus specificity by calculating the proportion of nonevent observations that were predicted to have an event response. ROC is the curve obtained by plotting sensitivity against one minus specificity. The ROC curve passes through the points (0, 0) and (1, 1) and lift up right. The area below the curve is one when the model discriminates perfectly, whereas the ROC curve lines from (0, 0) to (1, 1) when the model has no discriminating ability. In SAS the area below the ROC curve is given out and labeled as "c" under the title "Association of Predicted Probabilities and Observed Responses". From the plot of our models ROC curve we see that the curve is moderately satisfying and the statistic c is calculated as 0.613. This indicates that our models are reasonably useful and asthma prevalence rate can be explained by variation of environmental factors to some extent.

From the maximum likelihood analysis we see that for the ten provinces only the influence of Newfoundland, New Brunswick and Alberta are significant. These three provinces all have asthma prevalence lower than the average asthma prevalence and the negative value of their parameters indicates that living in these three provinces have a positive influence on people's health. People living in rural areas (GE36GU1) have a lower asthma prevalence. Adult with acceptable body mass (HWC6G2) tend to have less asthma prevalence than those with insufficient or some excess body mass. The parameters related to occupational variables indicate that the occupations of manufacturing non durable goods (LFC3), transportation (LFC6), finance (LFC9), community services (LFC10), personal services (LFC11), business services (LFC12) and public administration (LFC13) have a significantly adverse effect on people's health concerning asthma. People who engage in these occupations have a higher asthma prevalence. The estimate value of PAC6DPA1, PAC6DPA1 and PAC6DPA3 convey the relationship of physical activities with asthma. People who have less physical activities tend to get less asthma. This is consistent with our previous results and some findings in the asthma studies. Though physical activities do good to people, they also may increase the asthma prevalence. Relative research on asthma also indicates that there are many cases of exercise-induced asthma (see Manning, 1993, Tan et al. 2002 and website on Asthma and Allergy Information and Research (AAIR)). People born in Asia have much

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less asthma prevalence according to the large negative value of SDC6G3, while people born in Canada have the highest asthma prevalence over those born in Asia, Europe and USA. People who start smoking early (SMC6_31) have a higher asthma prevalence. The results also indicate that people who do not own the dwellings they live in (DHC6_02), who are female (DHC6_S2) or who are young (DHC6GAGE) have high asthma prevalence. The asthma prevalence for people who are married or have common-law or partner is smaller than those who are single (DHC6GMA2), widowed, separated or divorced (DHC6GMA3). Parameter estimates for variables representing household type and living arrangement (DHC6GE1, DHC6GE 4, DHC6GE5, DHC6GE 6, DHC6DL5, DHC6DL6) all show that those households in which parents are living with children have higher asthma prevalence.

From the results of the logistic regression we conclude that environmental factors do have a significant influence on asthma prevalence. It can to some extent explain the asthma distribution in the population.

4.5 Logistic Regression Analysis for Infants, Children and Adults

A possible difficulty with regression models fitted to the entire data set is that the relationships between response and predictors assumed by the models vary with age. In an attempt to examine this problem, we will fit separate regression models to the following age groups: infants (<=5 years), children (6-19 years) and adults (>=20 years). The sizes of these groups appear in Table 4.6. Note that the sizes of each group do not included observations with missing values.

7	A (A	•
Table	4 h	Age-group	CIZEC
1 avic	T •V	nec-eroup	SILCO.

	Infants	Children	Adults	total
Number of subjects	3,107	9,675	56,536	69,318

The logistic regression model was fitted to each age group following the approach described in Section 4.4 for the entire data, the results are reported in Table 4.7. ROC curves were also plotted in Figure 4.2.

Table 4.7 Stepwise logistic regression for infants, children and adults.

• Stepwise Logistic Regression Result for Infants.

Model Convergence Status Convergence criterion (GCONV=1E-8) satisfied. Model Fit Statistics Intercept Intercept and Criterion Only Covariates AIC 2576.873 2367.838 2582.914 2428.252 SC -2 Log L 2574.873 2347.838 Testing Global Null Hypothesis: BETA=0 Chi-Square DF Pr > ChiSq Test Likelihood Ratio 227.0348 9 <.0001 -9 236.3920 <.0001 Score Wald 198.1016 9 <.0001 Residual Chi-Square Test DF Pr > ChiSq Chi-Square 30.0237 21 0.0915 Analysis of Maximum Likelihood Estimates Standard Parameter \mathbf{DF} Estimate Error Chi-Square Pr > ChiSq 129.4231 -3.8072 Intercept 1 0.3347 <.0001 0.7219 -1.8112 SASK 1 6.2944 0.0121 25.5508 DHC6_02 1 0.6451 0.1276 <.0001 -0.9299 62.6915 <.0001 DHC6_S2 1 0.1174 1.1654 4.5139 1 1 DHC6GE5 0.2451 22.6065 <.0001 26.7407 DHC6GE9 0.8729 <.0001 1 1 1 1.6858 DHC6DL6 0.2248 56.2591 <.0001 30.0996 <.0001 DHC6DL9 0.9387 0.1711 0.0722 HWC6GHT 0.0112 41.7066 <.0001 <.0001 DHC6GAGE -0.8514 0.1378 38.1976 Association of Predicted Probabilities and Observed Responses Percent Concordant 63.0 Somers' D 0.279 0.284 Percent Discordant 35.1 Gamma 1.8 Tau-a 0.034 Percent Tied 581400 0.640 Pairs С Hosmer and Lemeshow Goodness-of-Fit Test Chi-Square DF Pr > ChiSq 49.4026 8 <.0001 8

• Stepwise Logistic Regression Result for Children

Model Convergence Status Convergence criterion (GCONV=1E-8) satisfied.

Model Fit Statistics

		intercept
	Intercept	and
Criterion	Only	Covariates
AIC	11392.583	10902.297
SC	11399.760	11081.730
-2 Log L	11390.583	10852.297

Testing Glob	oal Null Hypothe	esis: BETA	A=0
Test	Chi-Square	DF	Pr > ChiSq
Likelihood Ratio	538.2855	24	<.0001
Score	530.4603	24	<.0001
Wald	408.0939	24	<.0001

Residual	Chi-Squa	re Test
Chi-Square	DF	Pr > ChiSq
68.1231	33	0.0003

Analysis of Maximum Likelihood Estimates Standard

Standard					
Parameter	DF	Estimate	Error	Chi-Square	Pr > ChiSq
Intercept	1	-4.5118	0.4270	111.6541	<.0001
QUE	1	0.1511	0.0612	6.1009	0.0135
BC	1	0.2641	0.0746	12.5286	0.0004
GE36GU1	1	-0.2246	0.0833	7.2652	0.0070
LFC3	1	-0.6772	0.3038	4.9697	0.0258
LFC4	1	-1.3616	0.4711	8.3541	0.0038
LFC5	1	-0.7868	0.2942	7.1534	0.0075
lfC6	1	1.3229	0.2690	24.1775	<.0001
LFC7	1	0.8335	0.2450	11.5792	0.0007
LFC8	1	-0.3738	0.1308	8.1707	0.0043
lfC9	1	2.2393	0.2694	69.1067	<.0001
PAC6DPA2	1	0.3252	0.0712	20.8345	<.0001
SDC6G1	1	3.0832	0.3845	64.3091	<.0001
SDC6G2	1	1.9178	0.4723	16.4867	<.0001
DHC6_02	1	0.3432	0.0645	28.3153	<.0001
DHC6GB1	1	0.0845	0.0329	6.5891	0.0103
DHC6_S2	1	-0.1402	0.0514	7.4297	0.0064
DHC6GE3	1	0.9830	0.2366	17.2568	<.0001
DHC6GE6	1	1.4646	0.4027	13.2305	0.0003
DHC6DL2	1	0.6553	0.2394	7.4907	0.0062
DHC6DL5	1	1.6181	0.4924	10.8005	0.0010
DHC6DL6	1	0.2610	0.1178	4.9120	0.0267
DHC6DL8	1	0.3558	0.0864	16.9744	<.0001
HWC6GHT	1	-0.0399	0.00572	48.7487	<.0001
DHC6GAGE	1	0.2536	0.0426	35.3919	<.0001
Associa	tion of	Predicted	Probabilities	and Observed	Responses

Association	of	Predicted	Probabilities	and	Obs	served	Respon	ises
Percent	Cor	ncordant	55.0	Some	s'	D (D. 1 15	
Percent	Dis	scordant	43.5	Gamma	1	(0.117	
Percent	Tie	ed	1.5	Tau-a	ı	(0.025	
Pairs			10126314	с		(0.557	

Hosmer and Lemeshow Goodness-of-Fit Test Chi-Square DF Pr > ChiSq 92.4581 8 <.0001

• Stepwise Logistic Regression Result for Adults.

Model Convergence Status Convergence criterion (GCONV=1E-8) satisfied.

Model Fit Statistics

Test	AIC SC -2 Tes	Log L ting Global	Intercept Only 25049.257 25058.200 25047.257 Null Hypotl hi-Square	Intercept and Covariates 24257.856 24481.422 24207.856 nesis: BETA=0 DF Pr	> ChiSq
	lihood	Ratio	839.4011	24	<.0001
Scor	-		797.5723	24	<.0001
Wald			763.6152	24	<.0001
		Residua	1 Chi-Square	e Test	
	C	hi-Square	DF	Pr > ChiSq	
		114.9018	38	<.0001	
	Anal	ysis of Max	imum Likeli Standard	nood Estimates	
Parameter	DF	Estimate	Error	Chi-Square	Pr > ChiSq
Intercept	1	-0.9860	0.4695	4.4101	0.0357
NFLD	1	-0.6575	0.1705	14.8646	0.0001
NB	1	-0.3329	0.1271	6.8622	0.0088
QUE	1	-0.1011	0.0485	4.3480	0.0371
ONT	1	0.1385	0.0433	10.2546	0.0014
HWC6GB1	1	0.00921	0.00226	16.6009	<.0001
HWC6G2	1	-0.1969	0.0399	24.3546	<.0001
LFC1	1	0.3722	0.0493	57.0487	<.0001
LFC2	1	-0.4668	0.1915	5.9446	0.0148
LFC7	1	-0.4472	0.1403	10.1591	0.0014
LFC9	1	0.2247	0.0914	6.0391	0.0140
LFC10	1	0.2411	0.0550	19.2338	<.0001
PAC6DPA1	1	0.1412	0.0452	9.7538	0.0018
SDC6G1	1	0.8223	0.1411	33.9749	<.0001
SDC6G2	1	0.7091	0.1502	22.2745	<.0001
SDC6G3	1	-0.7501	0.2015	13.8527	0.0002
DHC6_02	1	0.2554	0.0455	31.5535	<.0001
DHC6GB1	1	-0.0506	0.0210	5.8018	0.0160
DHC6_S2	1	0.1652	0.0543	9.2641	0.0023
DHC6GM3	1	0.2135	0.0529	16.2800	<.0001
DHC6GE4	1	0.2751	0.0836	10.8310	0.0010
DHC6GE5	1	0.2647	0.0819	10.4438	0.0012
DHC6DL4	1	-0.2465	0.0443	30.8918	<.0001
HWC6GHT	1	-0.0337	0.00692	23.7150	<.0001
DHC6GAGE	1	-0.0776	0.00802	93.5061	<.0001

Association	of Predicted	Probabilities	and Observed	Responses
Percent	Concordant	60.4	Somers' D	0.227
Percent	Discordant	37.8	Gamma	0.231
Percent	Tied	1.8	Tau-a	0.029
Pairs		204207724	с	0.613

	Partition	for the Hosm	er and Leme	show Test	
		CCC6_1C	= YES	CCC6_10	2 = NO
Group	Total	Observed	Expected	Observed	Expected
1	5622	211	141.37	5411	5480.63
2	5688	262	219.57	5426	5468.43
3	5623	267	258.78	5356	5364.22
4	5619	305	296.30	5314	5322.70
5	5642	341	335.99	5301	5306.01
6	5661	340	377.48	5321	5283.52
7	5650	404	421.88	5246	5228.12
8	5668	440	480.04	5228	5187.96
9	5653	544	559.92	5109	5093.08
10	5710	764	778.68	4946	4931.32
	Hosmer ar Chi-So	nd Lemeshow G Tuare D			
		-	· · · ·		

53.6058 8 <.0001

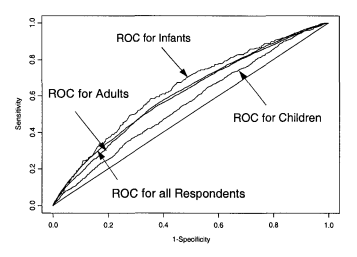


Figure 4.2 ROC curve for infants, children, adults and all respondents

From the results we see that the log likelihood ratio chi-square statistics for stepwise logistic regression models of infants, children and adults are all large enough to justify the significance of the model. However, all three models still show some lack of fit by the Hosmer and Lemeshow tests. Stepwise logistic regression results of all three groups of respondents show that there are relations between asthma prevalence and outdoor environment (provinces they live in), indoor environment (whether the dwelling they living is owned by family member, their living arrangement, etc.), and their basic physical characteristics (gender, height, age, etc.). The results for adults are similar to the results we get in section 4.4 for all respondents. However, they differ from those for infants and children. This may be explained by differences in the relationships across the groups but also by the fact that the adults group is the largest and thus this age group commands the trend in the entire data set. As would be expected, the ROC plot shown in Figure 4.2 shows that the ROC curve for adults and all respondents look very similar. The ROC curve for infants is relatively higher and the ROC curve for children is very low. These results indicate that for infants and adults environmental factors contained in our data set have moderate predictive power on asthma prevalence, but for children the predictive power is relatively low. This is an interesting point that deserves further research. We suspect that this may be caused by the fact that most children will spend much of their time in schools. Leickly (2003) estimated that people spend anywhere from 30 to 60% of their time in homes with the majority of the remaining time in an enclosed work area, or if it is a child, an indoor school environment. So other variables that describe school environment could be important to the study of the relation between children's asthma prevalence and environment.

4.6 Logistic Regression Model Validation

To acess the reliability of the logistic regression model, some limited cross validation was conducted. The subjects within each age group were randomly split into three subgroups using a trinomial distribution with equal probabilities (1/3, 1/3, 1/3). The first subgroup (Part A) was used for variable selection through stepwise logistic regression. The second subgroup (Part B) was used to fit the logistic regression model with the variables chosen in Part A. the third subgroup (Part C) is used to test the model. Figure 4.3 displays the ROC curves obtained from the test data (Part C) along with the ROCs for the groups from Figure 4.2.

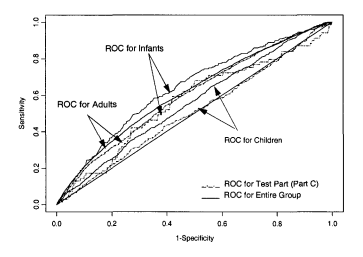


Figure 4.3 Cross validation ROC curve for infants, children and adults.

ROC curves of adults for entire group and test part are very similar to each other. This indicates the reliability of the results for adults. The corresponding ROC curves for infants and children in the entire groups and test parts differ from each other. However the ROC curves of infants are still high. Roc curves for children are all low, especially the ROC for the test part. From these results we conclude that the model fit results for infants and adults are reliable but that is not the case for children. Other factors may be need to study children's asthma prevalence as far as the logistic regression approach is concerned.

Chapter 5: Conclusions

The studies show that asthma rates are significantly different among provinces. The provinces of Nova Scotia and Newfoundland have the highest and lowest asthma rates, respectively. Compared to people living in urban areas, people living in rural areas have a significantly lower asthma prevalence. The asthma rate is also significantly different among the 26 or 33 regions divided by health area of the three provinces Ontario, Manitoba and Alberta. The overall message is that the regional outdoor environment influences the asthma prevalence rate.

Working environment is another factor that influences asthma prevalence and relative risk. Compared to people who have an occupation those who do not work have a higher asthma rate. For people who engage in different occupation the asthma risk they face varies significantly. People engaged in agriculture have the smallest asthma rate among all occupations. No matter whether we use the 21 occupations or the 13 occupations generated by the different classifications, people who engage in occupations related to service have relative higher asthma prevalence. Our study shows that finance, community services have an adverse effect on asthma.

Physical exercise influences the air change amount of our respiratory system. Iatrical studies find so many cases with exercise-induced asthma. Our studies also show that there is some relationship between the amount of physical exercise and asthma. Studies of our data which come from the NPHS survey of Canada suggest that people who have

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more physical exercise tend to have relatively higher asthma prevalence than others. Many cases with exercise-induced asthma have their first asthma attack during heavy exercise. It is deemed that the large amount and sudden inhale of cold and dry air trigger the symptoms of asthma. Warm up before exercise can lessen the stimuli to airway. Recent paper of Tan and Spector (2002) also indicate the benefits of warm up. So some exercise induced asthma can be prevented or lessen the asthma symptoms by not exercising so much and warming up sufficiently before exercising.

Smoking is another factor we consider in this study. Our study results show that for those who smoke the age people start smoking has very significant influence on their asthma prevalence. The earlier they start smoking, the higher the asthma prevalence among them. But among smokers the number of cigarettes they consume daily has no significant influence on asthma prevalence.

Our results show that demographic and social factors are another aspect we should consider when dealing with asthma. People who were born in Canada, America and Europe and Asia have asthma prevalence from the highest to the lowest one. Whether one has a well-balanced family structure can also have a significant influence on asthma rate. People who are married or have common-law or partner have a lower asthma prevalence compare to those who are single, widowed, separated or divorced.

Indoor environment can be another factor influencing asthma. Our study shows that the number of bedrooms in the dwelling, whether the dwelling is owned by household member, the household type and living arrangement all have an influence on asthma prevalence. Those people who live in dwellings owned by their family members, the dwelling they living in have more than two bedrooms have a less asthma prevalence.

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The study of the respondents' basic characteristics shows that gender is an influential factor for asthma. From the results of our analysis, we found that females have higher asthma prevalence than males. Children normally have higher asthma prevalence than adults. Adults with weight in the middle range have a lower asthma prevalence than those who are underweight or overweight.

Considering the above results we see that regional outdoor environment, working environment, physical exercise and air exchange amount, demographic and social environment, indoor environment and smoking pattern all affect asthma prevalence. They can to some extend explain the asthma distribution in the population.

These findings are consistent with results reported by other researchers, including Lierl and Hornung (2003), Maffei *et al.* (2001), FitzGerald *et al.* (2001), Ree and Kanabar (2000) and Tan and Spector (2002). Note that the data we used in this project is a cross sectional one. That is, the data are not collected prospectively. Thus some confounding may occur due to the fact that asthma may make people change their environment and behavior.

Our study is based on the Health File of the Public Use Microdata Documentation from the 1996-1997 National Population Health Survey. The Health File includes household residents in all provinces with the exclusion of populations on Indian Reserves, Canadian Forces Bases and some remote areas in Québec and Ontario. Accordingly, the findings in this project are applicable only to the scope of the Health File.

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APPENDIX A

Details of variable meaning and code for logistic regression analysis

(From the documentation for the NPHS provided by Statistics Canada)

		FORMATTED VALUE; DATA CLEAR; NEW NAME				
Variable Name	PRC6_CUR	Length	2	Position 7 -	8	
Question Name						$(0 \ 0 \ \cdots \ 0)$
Concept	Province of residence					$\begin{bmatrix} 0 & 0 & \cdots & 0 \\ 1 & 0 & \cdots & 0 \end{bmatrix}$
Question						
Universe	All respondents					
Note	Province of residence	at the time of data coll	lection in 199	6-1997.		$\left \begin{array}{cccc} \vdots & \ddots & 0\\ 0 & 0 & \cdots & 1 \end{array}\right $
Content			Code	Sample	Population	
NEWFOUNDLAND			10	963	549,322	PEI, NS, NB,
PRINCE EDWARD	ISLAND		11	918	132,322	QUE, ONT,
NOVA SCOTIA			12	986	895,914	MA, SASK,
NEW BRUNSWICK	č		13	1,032	728,118	ALB, BC
QUÉBEC			24	2,788	7,047,528	
ONTARIO			35	39,394	10,839,724	
MANITOBA			46	14,828	1,085,635	
SASKATCHEWAN			47	1,047	948,511	
ALBERTA			48	18,305	2,728,383	
BRITISH COLUMB	IA		59	1,543	3,686,279	
		Total		81,804	28,641,736	
Variable Name	GE36GURB	Length	1	Position 9		$\begin{pmatrix} 0 & 0 \\ 1 & 0 \end{pmatrix}$
Question Name	GENGUNB	Length	I	Position 9		
Concept	Derived rural and urbai	area - arouned				$\begin{pmatrix} 0 & 1 \end{pmatrix}$
Question	Denved foral and arbai	raiea - grouped				GE36GU1~2
Universe	All respondents					GE30GU1~2
Note	For sub-provincial anal			ta, use GE36GHRO and i ation on derived variables		
Content			<u>Code</u>	Sample	Population	
RURAL			1	2,659	2,888,000	
URBAN			2	4,903	6,105,200	
N/A - ON., MB., AB	., VAN. & MTL. CMA		6	74,242	19,648,535	
		Total		81,804	28,641,735	

	C	ORIGINAL V	ARIABI	Æ		FORMATTED VALUE DATA CLEAR NEW NAME
Variable Name	GE36GCMA	Length	3	Position 10	- 12	
Question Name						
Concept	Derived 1991 Census M	etropolitan Area - gr	ouped			
Question						
Universe	Respondents living in M					
Note	See documentation on o	lerived variables.				
Content			Code	Sample	Population	
MONTRÉAL			462	1,035	3,180,258	
VANCOUVER			933	680	1,814,536	
NOT APPLICABLE			996	80,089	23,646,941	
		Total		81,804	28,641,735	
Variable Name	DHC6_OWN	Length	1	Position	21	(0)
Question Name	HHLD_Q5					
Concept	Dwelling owned by house	ehold member				
Question	Is this dwelling owned by	a member of this he	ousehold (ev	en if being paid for)?		DHC6_OWN<=2
Universe	All respondents					_
Note						DHC61~2
<u>Content</u>			Code	Sample	Population	
YES			1	59,492	20,808,158	
NO			2	21,632	7,639,876	
DON'T KNOW			7	75	25,237	
REFUSAL			8	584	158,480	
NOT STATED			9	21	9,985	
		Total		81,804	28,641,736	
Variable Name	DHC6GBED	Length	2	Position 22	- 23	(0)
Question Name						
Concept	Derived number of bedro	ooms in dwelling - gr	ouped			
Question						
Universe	All respondents					
Note	Based on DHC6_BED.					$\begin{pmatrix} \cdot \\ 5 \end{pmatrix}$
Content			Code	Sample	Population	
NO BEDROOMS			0	326	116,393	DHC6GBED<=5
1 BEDROOM			1	7,217	1,919,568	
2 BEDROOMS			2	17,176	5,417,901	DHC6GB1
3 BEDROOMS			3	36,206	12,928,301	
4 BEDROOMS			4	14,961	6,003,835	
5 BEDROOMS OR I	MORE		5	4,335	1,866,890	
NOT STATED			99	1,583	388,847	
		Total		81,804	28,641,735	

	ORIGINAL VARIABLE						
Variable Name	GE36GHLR	Length	4	Position 13 - 16		NEW NAMI	
Question Name						/	
Concept	Derived health areas -	26 groups - grouped					
Question						-	
Universe	Respondents in Ontar	io, Manitoba and Alberta					
Note	See documentation or	n derived variables.					
Content			Code	Sample	Population		
NORTH AND SOUT	TH EASTMAN		461	2,953	86,530		
BURNTWOOD, NO	RMAN, PARKLAND		462	3,059	78,881		
CENTRAL, INTERL	AKE		463	2,803	153,601		
SOUTH WESTMAN	N, BRANDON, MARQUE	TTE	464	4,050	111,449		
WINNIPEG			465	1,963	655,174		
NORTHERN ALBE	RTA		481	4,488	418,276		
SOUTHERN ALBE	RTA		482	3,070	335,724		
CENTRAL ALBERT	ΓA		483	3,927	427,650		
CALGARY			484	3,694	842,824		
EDMONTON			485	3,126	703,909		
OTT-CAR.,PRES-F	RUS.,STOR.,DUN.,GLEN	RENF	3511	3,824	1,010,980		
LAN.,LE.,GREN.,H	AST.,P.E.,FRON.,LEN.,A	DD	3512	1,978	486,629		
N'UMBERLAND, V	ICT., HALIB.,PETERB., I	UR.	3521	3,131	759,832		
PEEL, HALTON			3522	3,221	1,243,008		
METRO TORONTO	o l		3523	4,085	2,238,057		
YORK, SIMCOE			3524	3,224	947,038		
NIAGARA			3531	1,506	406,163		
HAMILTON-WENT	WORTH		3532	1,631	484,221		
BRANT, HALDIMA	N, NORFOLK		3533	1,534	205,422		
WELLINGTON, DL	JFFERIN, WATERLOO		3534	3,101	634,539		
ESSEX, LAMBTON	I, KENT		3541	3,032	590,194		
ELGIN, MIDDLESE	EX, OXFORD		3542	1,688	609,183		
BRUCE, GREY, PE	ERTH, HURON		3543	1,573	297,485		
ALGOMA, COCHR	ANE, MANITOULIN, SU	DBURY	3551	2,952	443,511		
TIMISKAM., MUSK	OKA, PARRY SND., NIF	ISS.	3552	1,465	233,708		
THUNDER BAY, K	ENORA, RAINY RIVER		3561	1,449	249,755		
NOT APPLICABLE	:		9996	9,277	13,987,994		
		Total		81,804	28,641,736		

ORIGINAL VARIABLE								
Variable Name	GE36GHRO	Length	4	Position	17 - 20		NEW NAME	
Question Name								
Concept	Derived health areas -	33 groups - grouped						
Question								
Universe	Respondents in Ontar	io, Manitoba and Albe	rta					
Note	These health areas, c to official provincial he					s, do not correspor		
Content			Code	<u>San</u>	nple	Population		
NORTH AND SOU	TH EASTMAN		461	2,9)53	86,530		
BURNTWOOD, NO	ORMAN, PARKLAND		462	3,0)59	78,881		
CENTRAL, INTER	LAKE		463	2,8	303	153,601		
SOUTH WESTMA	N, BRANDON, MARQUE	TTE	464	4,0	50	111,449		
WINNIPEG			465	1,\$	963	655,174		
NORTHERN ALBE	RTA		481	4,4	188	418,276		
SOUTHERN ALBE	RTA		482	3,0	070	335,724		
CENTRAL ALBER	ТА		483	3,9	927	427,650		
CALGARY			484	3,6	694	842,824		
EDMONTON			485	3,1	126	703,909		
OTTAWA CARLET	FON		3511	2,6	350	738,276		
PRES-RUS.,STOF	R.,DUN.,GLEN.,RENF.		3512	1,1	174	272,703		
LAN, LE,GREN,HA	AST, P.E. FRONT, LENX,	ADD	3513	1,9	978	486,629		
N'UMBERLAND, V	ICT., HALIB., PETERB.		3521	1,8	513	277,441		
DURHAM			3522	1,0	618	482,391		
PEEL			3523	1,7	728	882,317		
METRO TORONT	0		3524	4,0	085	2,238,057		
YORK			3525	1,5	582	607,098		
SIMCOE			3526	1.0	342	339,940		
HALTON			3527	1,•	493	360,691		
NIAGARA			3531	1,	506	406,163		
HAMILTON-WENT	rworth		3532	1,6	531	484,221		
BRANT, HALDIMA	N, NORFOLK		3533	1,5	534	205,422		
WELLINGTON, D	UFFERIN		3534	1,	546	250,593	1	
WATERLOO			3536	1,\$	555	383,946		
ESSEX			3541	1,5	558	339,105		
LAMBTON, KENT			3542	1,-	474	251,089		
ELGIN, MIDDLESI	EX, OXFORD		3543	1,0	688	609,183		
BRUCE, GREY, P	ERTH, HURON		3544	1,	573	297,485		
ALGOMA, COCHF	RANE		3551	1,4	473	237,354		
MANITOULIN, SU	DBURY		3552	1.4	479	206,157	1	

	FORMATTED VALUE DATA CLEAR NEW NAME					
Variable Name	DHC6GAGE	Length	2	Position 37 -	38	(1)
Question Name						
Concept	Age - grouped					
Question						
Universe	All respondents					
Note	Based on DHC6_AGE.					(19)
Content			<u>Code</u>	Sample	Population	
0 TO 3 YEARS			1	2,452	807,493	(TREAT AS
4 TO 5 YEARS			2	1,355	860,745	CONTINUOUS
6 TO 9 YEARS			3	2,867	1,670,452	VARIABLE)
10 TO 11 YEARS			4	1,728	708,160	,
12 TO 14 YEARS			5	2,518	1,150,763	DHC6GAGE
15 TO 19 YEARS			6	4,449	2,110,798	
20 TO 24 YEARS			7	5,153	1,872,592	
25 TO 29 YEARS			8	6,362	2,008,407	
30 TO 34 YEARS			9	7,854	2,463,107	
35 TO 39 YEARS			10	8,021	2,739,289	
40 TO 44 YEARS			11	6,663	2,498,355	
45 TO 49 YEARS			12	5,642	2,044,499	
50 TO 54 YEARS			13	5,073	1,726,005	
55 TO 59 YEARS			14	4,313	1,402,356	
60 TO 64 YEARS			15	3,991	1,162,608	
65 TO 69 YEARS			16	3,993	1,133,537	
70 TO 74 YEARS			17	3,787	962,282	
75 TO 79 YEARS			18	2,719	681,003	
80 YEARS OR OLD)ER		19	2,864	639,287	
		Total		81,804	28,641,736	
Variable Name	DHC6_SEX	Length	1	Position	39	$\left(0\right)$
Question Name						$\left \left(1 \right) \right $
Concept	Sex					
Question						DHC6_S1~2
Universe	All respondents					
Note						
Content			<u>Code</u>	Sample	Population	
MALE			1	38,521	14,170,683	
FEMALE			2	43,283	14,471,052	
		Total		81,804	28,641,735	

ORIGINAL VARIABLE							FORMATT VALUE DATA CLE NEW NAM	
Variable Name Question Name	DHC6GMAR	Length	1	Position	40		$\begin{pmatrix} 0 & 0 \end{pmatrix}$	
Concept	Marital status - grouped							
Question	<i></i>						$\begin{pmatrix} 0 & 1 \end{pmatrix}$	
Universe	All respondents						DHC6GM	
Note	Based on DHC6_MAR.							
Content			Code	Sample	l	Population	DHC6GM	
MARRIED / COMM	ION-LAW / PARTNER		1	39,937		14,051,736		
SINGLE			2	28,616		11,561,153		
WIDOWED / SEPA	RATED / DIVORCED		3	13,090		2,998,280		
NOT STATED			9	161		30,566		
		Total		81,804		28,641,735		
Variable Name	DHC6GHSZ	Length	1	Position	41	·	(1)	
Question Name							2	
Concept	Derived household size - g	rouped						
Question								
Universe	All respondents						5	
Note	Based on DHC6DHSZ.							
<u>Content</u>			<u>Code</u>	Sample	2	Population	DHC6GH	
1 PERSON			1	16,028		3,171,363		
2 PERSONS			2	23,727		7,201,792		
3 PERSONS			3	14,358		5,391,788		
4 PERSONS			4	16,942		7,473,430		
5 OR MORE PERS	SONS	Total	5	10,749 81,804		5,403,362 28,641,735		
Variable Name	DHC6GLE5	Length	1	Position	42			
Question Name		-					$\begin{pmatrix} 1\\ 0 \end{pmatrix}$	
Concept	Derived persons <= 5 year	s old in household	- grouped					
Question							DHC6GL	
Universe	All respondents						Director	
Note	Based on DHC6_AGE.							
Content			Code	Sampl	<u>e</u>	Population		
Content			1	16,460		6,284,053		
YES			2	65,344		22,357,682		
			2					

	FORMATTED VALUE DATA CLEAR NEW NAME					
Variable Name	DHC6G611	Length	1	Position	43	(1)
Question Name						
Concept	Derived persons 6 to 11	years old in hhld - gr	rouped			
Question						DHC6G61
Universe	All respondents					
Note	Based on DHC6_AGE.					
<u>Content</u>			Code	Sample	Population	
YES			1	18,209	7,576,504	
NO			2	63,595	21,065,231	
		Total		81,804	28,641,735	
Variable Name	DHC6GECF	Length	2	Position 44	- 45	(1 0 0)
Question Name						
Concept	Derived household type	- grouped				
Question						
Universe	All respondents					$\left(\begin{array}{ccc} \cdot & \cdot & -1 \\ 0 & 0 & \cdots & 0 \end{array}\right)$
Note	Based on the relationsh	p matrix. See docum	entation on o	terived variables.		
Content			Code	Sample	Population	DHC6GECF<=9
COUPLE WITH CH	HILDREN < 25		1	31,062	13,239,101	
COUPLE W/WO C	HILDREN >= 25, W/WO O	THERS	2	2,772	1,247,518	
SINGLE			3	16,028	3,171,363	DHC6GE1~6,
SINGLE WITH OT	HERS		4	2,417	787,252	8~9
COUPLE WITH CH	HILDREN < 25, OTHERS		5	3,198	1,694,119	
COUPLE ALONE			6	17,967	5,384,483	
SINGLE PARENT,	CHILDREN < 25		7	5,535	2,068,694	
OTHER SINGLE P	ARENT HOUSEHOLD		8	2,212	840,530	
OTHER			9	597	204,347	
NOT STATED			99	16	4,330	
		Total		81,804	28,641,737	

		ORIGINAL V	ARIABLE			FORMATTED VALUE DATA CLEAR NEW NAME
Variable Name	DHC6DLVG	Length	2	Position 46 - 4	7	$(1 \ 0 \ \cdots \ 0)$
Question Name						0 1 0
Concept	Derived living arranger	nents of the selected i	respondent			0 0
Question						
Universe	All respondents					
Note	Based on the relations	hip matrix. See docum	entation on deri	ved variables.		
Content			<u>Code</u>	Sample	Population	DHC6DLVG<=
UNATTACHED IND	VIDUAL LIVING ALONE		1	16,028	3,171,363	10
UNATTACHED IND	VIDUAL LIVING WITH C	THERS	2	2,211	704,020	
LIVING WITH SPOU	JSE / PARTNER		3	17,966	5,384,406	DHC6DL1~6,
PARENT LIVING W	T SPOUSE/PARTNER,C	HILDREN	4	19,818	7,632,923	8~9
SINGLE PARENT L	VING WITH CHILDREN		5	3,566	1,043,680	
CHILD LIVING WITH	I SINGLE PARENT		6	1,525	604,825	
CHILD LIVING WT :	SINGLE PARENT, SIBLI	NGS	7	1,707	891,491	
CHILD LIVING WITI	H TWO PARENTS		8	3,157	1,085,852	
CHILD LIVING WITI	H TWO PARENTS, SIBL	INGS	9	10,174	5,434,898	
OTHER			10	5.640	2,685,570	
NOT STATED			99	12	2,707	
		Total		81,804	28,641,735	
Variable Name	CCC6_1C	Length	1	Position	165	Respond
Question Name	CHR-Q1					Variable
Concept	Has asthma					
Question	Do you have asthma o	diagnosed by a health	professional?			CCC6_1C<=2
Universe	All respondents					
Note						
Content			Code	Sample	Population	
YES			1	6,242	2,236,139	
NO			2	75,512	26,394,976	
DON'T KNOW			7	38	7,859	
REFUSAL			8	5	523	
NOT STATED			9	7	2,239	
		Total		81,804	28,641,736	

	FORMATTED VALUE DATA CLEAR NEW NAME					
Variable Name	CCC6_1H	Length	1	Position 1	84	
Question Name	CHR-Q1					
Concept	Has chronic bronchitis or	emphysema				
Question	Do you have chronic bror	ichitis or emphysem	na diagnosed	by a health professional?		
Universe	All respondents					
Note						
Content			<u>Code</u>	Sample	Population	
YES			1	2,603	835,012	
NO			2	79,147	27,795,358	
DON'T KNOW			7	31	6,912	
REFUSAL			8	5	1,235	
NOT STATED			g	18	3,219	
		Total		81,804	28,641,736	
Variable Name	SDC6GCB	Length	2	Position 207 - 2	08	
Question Name	~~~~~	renhm	é.,	FV91UUH &V(+ &	~~	
Concept	Country of birth - grouped	4				0 1 0
Question	country of pintit - grouper	1				0 0 1
	All environments					
Universe	All respondents					
Note	Based on SDC6_1.					SDC6GCB<=4
Content			<u>Code</u>	Sample	Population	
CANADA			1	69,628	23,771,682	SDCGG1~3
U.S., EUROPE, AU	STRALIA		2	8,353	2,663,564	
ASIA			3	2,203	1,418,177	
OTHER			4	1,479	737,741	
NOT STATED			99	141	50,571	
		Total		81,804	28,641,735	
Variable Name	SDC6GRAC	Length	2	Position 213 - 2	14	(1)
Question Name						
Concept	Derived race or colour - g	rouped				
Question						SDC6GRAC<=2
Universe	All respondents					
Note	Based on SDC6_7A to S	DC6_7L.				SDC6GR1
Content			<u>Code</u>	Sample	Population	
WHITE			1	75,423	25,461,656	
OTHER			2	6,022	3,079,856	
NOT STATED			99	359	100,224	
		Total		81,804	28,641,736	1

	FORMATTED VALUE DATA CLEAR NEW NAME						
Variable Name	HWC6G3KG	Length	4	Position	290 • 29	3	$\left(3\right)$
Question Name							4
Concept	Weight in kilograms - g	rouped					
Question							137
Universe	All respondents						
Note	Based on HWC6_3KG. 113 kgs. Males aged 2 who were < 40 kgs wer	0 & over who were < 5					HWC6G3KG <=137
Content			<u>Code</u>	Sam	nple	Population	
KILOGRAMS			3 - 137	78,3	43	27,690,651	HWC6G3KG
NOT STATED			999	3,4	161	951,084	
		Total		81,8	304	28,641,735	
Variable Name	LFC6GI13	Length	2	Position	227 .	228	
Question Name							$\begin{pmatrix} 1 & 0 & \cdots & 0 \\ 0 & 1 & \cdots & 0 \end{pmatrix}$
Concept	Industry Codes for ma	ain job - 13 groups - gr	ouned				
Question	madolity coase ka me	ninjob togroapo gr	oupou				0 0
Universe	Respondents who ans						
Note	Based on LFC6CSIC.						
	based on Li Obcolo.						
<u>Content</u>			<u>Code</u>		ample	Population	LFC6GI13<=96
AGRICULTURAL			1		2,124	404,842	LFC1~13
OTHER PRIMARY			2		1,456	362,431	LFC1~13
MANUFACTURING	/NON-DURABLE		3		2,580	1,097,129	
MANUFACTURING	/ DURABLE		4		2,815	991,389	
CONSTRUCTION			5		2,494	831,437	
TRANSPORTATION	N		6		3,310	1,133,835	
WHOLESALE TRAI	DE		7		1,652	616,504	
RETAIL TRADE			8		5,125	1,890,903	
FINANCE			9		2,023	775,239	
COMMUNITY SERV	/ICES		10		8,407	2,732,158	
PERSONAL SERVI	CES		11		3,912	1,524,934	
	ES		12		3,975	1,579,035	
BUSINESS SERVIC	RATION		13		2,546	924,500	
BUSINESS SERVIC PUBLIC ADMINIST			96	3	6,292	12,927,598	
					3,093	849,800	
PUBLIC ADMINIST			99			045,000	

ORIGINAL VARIABLE								
Variable Name	LFC6GO21	Length	2	Position 225 - 2	26	NEW NAME		
Question Name								
Concept	Occupation Codes fo	r main job - 21 groups ·	grouped					
Question								
Universe	Respondents who ar	iswered LFC6_2=1						
Note	Based on LFC6CSO	C. See documentation (on derived var	iables.				
Content			<u>Code</u>	Sample	Population			
MANAGERIAL, AD	MINISTRATIVE		1	5,825	2,062,784			
NATURAL SCIENC	E		2	1,780	601,459			
SOCIAL SCIENCE			3	994	358,871			
RELIGION			4	121	26,294			
TEACHING			5	2,232	740,580			
MEDICINE			6	2,338	739,736			
ARTISTIC			7	868	319,723			
CLERICAL			8	5,694	2,243,093			
SALES			9	3,796	1,365,650			
SERVICES			10	5,733	2,171,099			
FARMING			11	2,111	418,617			
FISHING			12	96	53,532			
FORESTRY			13	188	85,416			
MINING			14	408	53,215			
PROCESSING			15	1,011	383,650			
MACHINING			16	696	191,581			
FABRICATING			17	3,004	1,149,963			
CONSTRUCTION			18	2,422	800,219			
TRANSPORTATIO	N		19	1,770	564,030			
MATERIALS HANE	LING		20	888	393,366			
OTHER CRAFTS			21	444	141,460			
NOT APPLICABLE			96	36,292	12,927,598			
NOT STATED			99	3,093	849,800			
		Total		81,804	28,641,735			

		ORIGINAL VA	ARIABL	E		·	FORMATTED VALUE DATA CLEAR NEW NAME	
Variable Name Question Name Concept Question Universe Note Content BMI SCORE NOT APPLICABLE NOT STATED	Respondents aged 2	Length ndex (1 decimal place) -) to 64 years old excludi , HWC6G3KG and HW 1	ing pregnant	documentation	296 - 2 n on derived v <u>Samble</u> 50,347 29,474 1,983		(13.2 ⋮ 57.9 0 HWC6GBMI<=99.6 HWC6GB1	
Variable Name Question Name Concept Question Universe) to 64 years old excludi		Position	81,804 3	28,641,736	$ \left \begin{array}{cccccccccccccccccccccccccccccccccccc$	
Note <u>Content</u> INSUFFICIENT WEI ACCEPTABLE WEI SOME EXCESS WE OVERWEIGHT NOT APPLICABLE NOT STATED	іднт GHT	11. See documentation o Total	on derived va <u>Code</u> 1 2 3 4 6 9	<u>5</u>	Sample 3,805 21,014 9,779 15,749 29,474 1,983 81,804	Population 1,422,385 7,474,777 3,287,154 4,980,393 10,939,358 537,668 28,641,735	HWC6GSW<=6	

	FORMATTED VALUE DATA CLEAR NEW NAME					
Variable Name	HWC6GHT	Length	2	Position 288 - 2	289	
Question Name						$\left(\begin{array}{c} 2 \end{array} \right)$
Concept	Height - adults and child	dren - grouped				
Question						
Universe	All respondents					64
Note				. Females > 5*11" were gi ales aged 20 + who were		HWC6GHT<=64
<u>Content</u>			<u>Code</u>	Sample	Population	
1'0" (12 INCHES) (29.2 TO 31.7 CM)		2	1	113	
1'2" (14 INCHES) (34.3 TO 36.7 CM)		4	1	160	
1'3" (15 INCHES) (3	36.8 TO 39.3 CM)		5	2	613	
1'4" (16 INCHES) (39.4 TO 41.8 CM)		6	3	490	
1'5" (17 INCHES) (4	41.9 TO 44.4 CM)		7	2	234	
1'6" (18 INCHES) (44.5 TO 46.9 CM)		8	10	4,346	
1'7" (19 INCHES) (4	47.0 TO 49.4 CM)		9	4	306	
1'8" (20 INCHES) (-	49.5 TO 52.0 CM)		10	16	1,588	
1'9" (21 INCHES) (52.1 TO 54.5 CM)		11	26	2,781	
1'10" (22 INCHES)	(54.6 TO 57.1 CM)		12	39	3,622	
1'11" (23 INCHES)	(57.2 TO 59.6 CM)		13	54	4,351	
2'0" (24 INCHES) (59.7 TO 62.1 CM)		14	133	36,978	
2'1" (25 INCHES) (62.2 TO 64.7 CM)		15	52	7,319	
2'2" (26 INCHES) (64.8 TO 67.2 CM)		16	84	10,392	
2'3" (27 INCHES) (67.3 TO 69.8 CM)		17	54	11,098	
2'4" (28 INCHES) (69.9 TO 72.3 CM)		18	81	14,271	
2'5" (29 INCHES) (72.4 TO 74.8 CM)		19	89	15,413	
2'6" (30 INCHES) (74.9 TO 77.4 CM)		20	208	67,043	
27" (31 INCHES) (77.5 TO 79.9 CM)		21	61	19,928	
2'8" (32 INCHES) (80.0 TO 82.5 CM)		22	101	19,085	
2'9" (33 INCHES) (82.6 TO 85.0 CM)		23	94	25,673	
2'10" (34 INCHES)	(85.1 TO 87.5 CM)		24	61	12,939	
2'11" (35 INCHES)	(87.6 TO 90.1 CM)		25	83	43,312	
3'0" (36 INCHES) (90.2 TO 92.6 CM)		26	643	214,957	
3'1" (37 INCHES) (92.7 TO 95.2 CM)		27	112	65,670	
3'2" (38 INCHES) (95.3 TO 97.7 CM)		28	154	100,600	
3'3" (39 INCHES) (97.8 TO 100.2 CM)		29	110	80,384	
	100.3 TO 102.8 CM)		30	199	129,059	
3'5" (41 INCHES) (102.9 TO 105.3 CM)		31	171	103,909	
	105.4 TO 107.9 CM)		32	391	255,468	
3'6" (42 INCHES) (115	96,566	1

ORIGINAL VARIABLE							
3'8" (44 INCHES) (110.5 TO 112.9 CM)	34	136	98,490	NEW NAME			
3'9" (45 INCHES) (113.0 TO 115.5 CM)	35	133	89,220				
3'10" (46 INCHES) (115.6 TO 118.0 CM)	36	97	68,852				
3'11" (47 INCHES) (118.1 TO 120.6 CM)	37	116	98,204				
4'0" (48 INCHES) (120.7 TO 123.1 CM)	38	1,026	576,944				
4'1" (49 INCHES) (123.2 TO 125.6 CM)	39	152	122,648				
"2" (50 INCHES) (125.7 TO 128.2 CM)	40	251	171,253				
'3" (51 INCHES) (128.3 TO 130.7 CM)	41	162	124,988				
'4" (52 INCHES) (130.8 TO 133.3 CM)	42	155	138,897				
'5" (53 INCHES) (133.4 TO 135.8 CM)	43	201	120,695				
1'6" (54 INCHES) (135.9 TO 138.3 CM)	44	339	191,507	l			
'7" (55 INCHES) (138.4 TO 140.9 CM)	45	218	103,259				
'8" (56 INCHES) (141.0 TO 143.4 CM)	46	298	136,943				
'9" (57 INCHES) (143.5 TO 146.0 CM)	47	260	115,720				
'10" (58 INCHES) (146.1 TO 148.5 CM)	48	503	195,802				
'11" (59 INCHES) (148.6 TO 151.0 CM)	49	825	336,618				
50" (60 INCHES) (151.1 TO 153.6 CM)	50	2,909	1,087,011				
51" (61 INCHES) (153.7 TO 156.1 CM)	51	2,748	954,010				
5'2" (62 INCHES) (156.2 TO 158.7 CM)	52	5,674	1,843,299				
5'3" (63 INCHES) (158.8 TO 161.2 CM)	53	5,249	1,847,610				
5'4" (64 INCHES) (161.3 TO 163.7 CM)	54	6,712	2,118,939				
5'5" (65 INCHES) (163.8 TO 166.3 CM)	55	5,566	1,952,506				
5'6" (66 INCHES) (166.4 TO 168.8 CM)	56	6,593	2,209,327				
57" (67 INCHES) (168.9 TO 171.4 CM)	57	6,193	2,090,665				
5'8" (68 INCHES) (171.5 TO 173.9 CM)	58	6,499	2,107,348				
5'9" (69 INCHES) (174.0 TO 176.4 CM)	59	4,576	1,574,484				
5'10" (70 INCHES) (176.5 TO 179.0 CM)	60	5,536	1,958,810				
"11" (71 INCHES) (179.1 TO 181.5 CM)	61	4,823	1,533,474				
50" (72 INCHES) (181.6 TO 184.1 CM)	62	4,194	1,294,319	1			
51" (73 INCHES) (184.2 TO 186.6 CM)	63	1,837	548,644				
5'2" (74 INCHES) (186,7 TO 189.1 CM)	64	2,864	918,287				
NOT STATED	99	1,805	564,298				
	Total	81,804	28,641,736				

Question Name 0 1 Concept Derived participant in leisure physical activity 0 Question 0 0			ORIGINAL V	ARIABL				FORMATTED VALUE DATA CLEAR NEW NAME		
ConceptDerived participant in leisure physical activityQuestionUniverseRespondents aged 12 and overNoteBased on PAC6_1A to PAC6_1X.PAC6DLEI<	/ariable Name	PAC6DLEI	Length	1	Position	796		$\begin{pmatrix} 1 & 0 \end{pmatrix}$		
Constant Code Sample Population Universe Respondents aged 12 and over PAC6DLEI<	Juestion Name							0 1		
Constant Code Sample PAC6DLEI< Variable Name PAC6DLE 6 8,402 4,046,849 PAC6DLEI Variable Name PAC6DEF 2 8,073 2,655,779 0.01 0.01 2 8,073 2,655,779 0.01 <td>Concept 201</td> <td>Derived participant in</td> <td>elsure physical activity</td> <td></td> <td></td> <td></td> <td></td> <td></td>	Concept 201	Derived participant in	elsure physical activity							
TACODUCING TACODUCING Content Code Sample Population TEXENTLE PAC6DL1 PAC6DL1 PAC6DL1 PAC6DL1 PAC6DL1 PAC6DL1 Total Bis04 PAC6DL1 Total Bis04 PAC6DL1 Variable Name PAC6DFM Length 3 Position 795 Concept Derived monthly freq; of physical adivity lasting > 15 min. Concept Population Note Sample Population Motiverse Respondents aged 12 and over Note Sample Population MOT HPPLICABLE 996 8,402 4,046,849 NOT STATED 999 2,025 7.95 Question Name	Juestion									
Content YES Code 1 Sample 63,304 Population 21,280,090 PAC6DL1 VTAPPLICABLE 6 8,402 4,046,849 A NOT APPLICABLE 6 8,402 4,046,849 A NOT STATED 9 2,025 759,016 A A Variable Name PAC6DFM Length 3 Position 797<-799	Jniverse	Respondents aged 12	and over					PAC6DLEI<=6		
Convert Sample Convert Convert <th< td=""><td>lote</td><td>Based on PAC6_1A to</td><td>PAC6_1X.</td><td></td><td></td><td></td><td></td><td></td></th<>	lote	Based on PAC6_1A to	PAC6_1X.							
NO 2 8,073 2,555,779 NOT APPLICABLE 6 8,402 4,046,849 NOT STATED 9 2,025 759,018 Total 81,804 28,641,736 Variable Name PAC6DFM Length 3 Position 797 7.99 Question Name Concept Derived monthly freq, of physical activity lasting > 15 min. 71 23,835,869 0 0 1 255 0 0 Moters Respondents aged 12 and over Note Based on PAC6_2A to PAC6_2X. See documentation on derived variables. PAC6DFM<	Content			Code	Sample	<u>1</u>	Population	PAC6DL1		
NOT APPLICABLE 6 8,402 4.046,849 NOT STATED 9 2,025 759,018 Total 81,804 28,641,736 Variable Name PAC60FM Length 3 Position 797 799 Question Name Concept Derived monthly freq. of physical addivity lasting > 15 min. 797 799 001 1 1 2255 0 0 Question Concept Derived monthly freq. of physical addivity lasting > 15 min. 9 9 2,025 759,018 0 0 0 1 1 255 0	/ES			1	63,304	-	21,280,090			
VOT STATED 9 2,025 759,018 Total 81,804 28,641,736 Variable Name PAC6DFM Length 3 Position 797 799 Question Name Derived monthly freq. of physical activity lasting > 15 min. 0.01 1 1 255 0 Question Based on PAC6_2A to PAC6_2X. See documentation on derived variables. Population PAC6DFM<	10			2	8,073		2,555,779			
NOT STATED 9 2,025 759,018 Total 81,804 28,641,736 Variable Name PAC6DFM Length 3 Position 797 799 Question Name Derived monthly freq, of physical activity lasting > 15 min. 797 799 (0,01) 1 255 0 0 Question Moverse Respondents aged 12 and over Variable science Science Science 255 0<	NOT APPLICABLE			6			4,046,849			
Total81,80428,641,736Variable NamePAC6DFMLength3Position797799Question NameConceptDerived monthly freq. of physical activity lasting > 15 min. (0.01) 1QuestionQuestionRespondents aged 12 and over (0.01) 1NoteBased on PAC6_2A to PAC6_2X. See documentation on derived variables.PAC6DFM<				9						
Variable Name PAC6DFM Length 3 Position 797 799 Question Name Concept Derived monthly freq, of physical activity lasting > 15 min. 0.01 1 255 0 Question Universe Respondents aged 12 and over 0 255 71,377 23,835,869 PAC6DFM<<=99			Total							
Question Name 1 Concept Derived monthly freq, of physical activity lasting > 15 min. Question 255 Universe Respondents aged 12 and over Note Based on PAC6_2A to PAC6_2X. See documentation on derived variables. Content Code Sample Population MONTHLY FREQUENCY 0 - 255 71,377 23,835,869 PAC6DFM<=99	Variable Name	PAC6DFM		3		. 799		(0.01)		
Concept Derived monthly freq. of physical activity lasting > 15 min. Image: Concept Concent Concept Concept Concent Concept Concept Concept	Question Name		·							
Universe Respondents aged 12 and over Action Pack (2,2,3) Pack (2,2,3) Pack (2,2,3) Pack (2,3,3)	Concept									
Universe Respondents aged 12 and over 0 Note Based on PAC6_2A to PAC6_2X. See documentation on derived variables. 0 Content Code Sample Population MONTHLY FREQUENCY 0 - 255 71,377 23,835,869 NOT APPLICABLE 996 8,402 4,046,849 NOT STATED 999 2,025 759,018 Variable Name PAC6DEE Length 4.1 Position 792 - 795 Question Name Concept Derived energy expenditure (1 decimal point) Variable Name Variable Name Variable Name Pac6DEE Variable Name Pac6det Variable Name Variable Name Pac6det Variable Name Variable Name Variable Name Pac6det Variable Name Variable Name Pac6det Variable Name Variable Name<	Question									
Note Based on PAC6_2A to PAC6_2X. See documentation on derived variables. PAC6DFM<=99 Content 0 - 255 71,377 23,835,869 MONTHLY FREQUENCY 0 - 255 71,377 23,835,869 NOT APPLICABLE 996 8,402 4,046,849 NOT STATED 999 2,025 759,018 Variable Name PAC6DEE Length 4.1 Position Variable Name PAC6DEE Length 4.1 Position 792 795 Question Name Concept Derived energy expenditure (1 decimal point) Variable Name Variable Name PAC6_1 to PAC6_1 to PAC6_2. See documentation on derived variables. Concept Derived energy expenditure (1 decimal point) Variable Name Population Universe Respondents aged 12 and over Sample Population Note Based on PAC6_1 to PAC6_1 to PAC6_2. See documentation on derived variables. Content Content Code Sample Population AMOUNT OF ENERGY EXPENDITURE 0.0 - 34.4 71,377 23,825,869 NOT APPLICABLE 99.6 8,402 4,046,849 NOT STATED 99.9 2,025 759,018	Universe	Respondents aged 12	and over							
ContentCodeSamplePopulationMONTHLY FREQUENCY0 - 25571,37723,835,869PAC6DFM<=99	Note	Based on PAC6 2A b								
MONTHLY FREQUENCY 0 - 255 71,377 23,835,869 PAC6DFM1 NOT APPLICABLE 996 8,402 4,046,849 PAC6DFM1 NOT STATED 999 2,025 759,018 Total 81,804 28,641,735 Variable Name PAC6DEE Length 4.1 Position 792 - 795 Question Name Concept Derived energy expenditure (1 decimal point) 792 - 795 Question Universe Respondents aged 12 and over Variable Add Over Variable Name	Content		PAC6DFM<=996							
NOT APPLICABLE 996 8,402 4,046,849 PAC6DFM1 NOT STATED 999 2,025 759,018 759,018 Total 81,804 28,641,735 28,641,735 Variable Name PAC6DEE Length 4.1 Position 792 - 795 Question Name Concept Derived energy expenditure (1 decimal point) 792 - 795 Question Name Universe Respondents aged 12 and over Variable S. Note Based on PAC6_1X and PAC6_2. See documentation on derived variables. Eontent Content Code Sample Population AMOUNT OF ENERGY EXPENDITURE 0.0 - 34.4 71,377 23,835,869 NOT APPLICABLE 99.6 8,402 4,046,849 NOT STATED 99.9 2,025 759,018		ENCY				-				
NOT STATED 999 2,025 759,018 Total 81,804 28,641,735 Variable Name PAC6DEE Length 4.1 Position 792 - 795 Question Name Concept Derived energy expenditure (1 decimal point) 792 - 795 Question Name Universe Respondents aged 12 and over Variable Ac6_1A to PAC6_1X and PAC6_2. See documentation on derived variables. Content Code Sample Ponulation AMOUNT OF ENERGY EXPENDITURE 0.0 - 34.4 71,377 23,835,869 NOT APPLICABLE 99.6 8,402 4,046,849 NOT STATED 99.9 2,025 759,018								PAC6DFM1		
Variable NamePAC6DEELength4.1Position792795Question NameConceptDerived energy expenditure (1 decimal point)QuestionUniverseRespondents aged 12 and overNoteBased on PAC6_1A to PAC6_1X and PAC6_2. See documentation on derived variables.ContentCodeSamoleSamoleAMOUNT OF ENERGY EXPENDITURE0.0 - 34.471,37723,835,869NOT APPLICABLE99.92,025759,018										
Concept Derived energy expenditure (1 decimal point) Question Universe Respondents aged 12 and over Note Based on PAC6_1A to PAC6_1X and PAC6_2. See documentation on derived variables. Content Code Sample Population AMOUNT OF ENERGY EXPENDITURE 0.0 - 34.4 71,377 23,835,869 NOT APPLICABLE 99.6 8,402 4,046,849 NOT STATED 99.9 2,025 759,018			Total							
Concept Derived energy expenditure (1 decimal point) Question Universe Respondents aged 12 and over Note Based on PAC6_1A to PAC6_1X and PAC6_2. See documentation on derived variables. Content Code Sample Population AMOUNT OF ENERGY EXPENDITURE 0.0 - 34.4 71,377 23,835,869 NOT APPLICABLE 99.6 8,402 4,046,849 NOT STATED 99.9 2,025 759,018										
ConceptDerived energy expenditure (1 decimal point)QuestionUniverseRespondents aged 12 and overNoteBased on PAC6_1A to PAC6_1X and PAC6_2. See documentation or derived variables.ContentCodeSampleAMOUNT OF ENEREXPENDITURE0.0 - 34.471,37723,835,869NOT APPLICABLE99.68,4024,046,849NOT STATED99.92,025759,018		PACODEE	Length	4.1	Position 792	2 - 795				
Question Respondents aged 12 and over Note Based on PAC6_1A to PAC6_1X and PAC6_2. See documentation on derived variables. Content Code Sample AMOUNT OF ENERGY EXPENDITURE 0.0 - 34.4 71,377 23,835,869 NOT APPLICABLE 99.6 8,402 4,046,849 NOT STATED 99.9 2,025 759,018		P 1								
Universe Respondents aged 12 and over Note Based on PAC6_1A to PAC6_1X and PAC6_2. See documentation on derived variables. Content Sample Population AMOUNT OF ENERGY EXPENDITURE 0.0 - 34.4 71,377 23,835,869 NOT APPLICABLE 99.6 8,402 4,046,849 NOT STATED 99.9 2,025 759,018	-	Derived energy exper	iditure (1 decimal point))						
Note Based on PAC6_1A to PAC6_1X and PAC6_2. See documentation on derived variables. Content Code Sample Population AMOUNT OF ENERGY EXPENDITURE 0.0 - 34.4 71,377 23,835,869 NOT APPLICABLE 99.6 8,402 4,046,849 NOT STATED 99.9 2,025 759,018		m () · · · ·								
Content Code Sample Population AMOUNT OF ENERGY EXPENDITURE 0.0 - 34.4 71,377 23,835,869 NOT APPLICABLE 99.6 8,402 4,046,849 NOT STATED 99.9 2,025 759,018										
AMOUNT OF ENERGY EXPENDITURE 0.0 - 34.4 71,377 23,835,869 NOT APPLICABLE 99.6 8,402 4,046,849 NOT STATED 99.9 2,025 759,018	Note	Based on PAC6_1A t	p PAC6_1X and PAC6	_2. See docu	imentation on derive	ı variables.				
NOT APPLICABLE99.68,4024,046,849NOT STATED99.92,025759,018	<u>Content</u>			<u>Code</u>	Sampl	2	Population			
NOT STATED 99.9 2,025 759,018	AMOUNT OF ENEF	rgy expenditure		0.0 - 34.4	71,377		23,835,869			
	NOT APPLICABLE			99.6	8,402		4,046,849			
Total 81,804 28,641,735				99.9	2,025		759,018			
	NOT STATED				01 004		20 6/1 725			

	ORIGINAL VARIABLE								
Variable Name	PAC6DFR	Length	1	Position	800				
Question Name									
Concept	Derived frequency of all phy	sical activity							
Question									
Universe	Respondents aged 12 and c								
Note	Based on PAC6DFM. See c	ocumentation on c	lerived varia	ables.					
Content			Code	Sample	P	opulation			
REGULAR			1	42,545	14	1,156,094			
OCCASIONAL			2	12,749	4	4,536,763			
INFREQUENT			3	16,083		5,143,012			
NOT APPLICABLE			6	8,402	4	4,046,849			
NOT STATED			9	2,025		759,018			
		Total		81,804		3,641,736			
Variable Name	PAC6DFD	Length	1	Position	801				
Question Name									
Concept	Derived participation / daily								
Question									
Universe	Respondents aged 12 and c	ver							
Note	Based on PAC6DFM. See (
<u>Content</u>			<u>Code</u>	Sample	<u>P</u>	opulation			
DAILY			1	22,601		7,305,908			
NOT DAILY			2	48,776	1	6,529,961			
NOT APPLICABLE			6	8,402		4,046,849			
NOT STATED			9	2,025		759,018			
		Total		81,804	2	8,641,736			
Variable Name	PAC6DPAI	Length	1	Position	802		$(1 \ 0 \ 0)$		
Question Name							0 1 0		
Concept	Derived physical activity inc	ex					0 0 1		
Question									
Universe	Respondents aged 12 and								
	Based on PAC6DEE. See (locumentation on (derived vari	ables.			PAC6DPAI<=		
Note			Code	Sample	E	Population			
Note <u>Content</u>			1	14,377		4,911,009			
						5 000 005			
Content			2	16,346		5,382,985			
Content ACTIVE				16,346 40,654		5,382,985 3,541,875			
<u>Content</u> ACTIVE MODERATE			2		1				
<u>Content</u> ACTIVE MODERATE INACTIVE			2 3	40,654	1	3,541,875			

	FORMATTED VALUE DATA CLEAR NEW NAME						
Variable Name Question Name Concept	SMC6_3 SMK-Q3 Age started smoking		3	Position	927 -	929	
luestion		begin to smoke cigarette	es daily?				85
Iniverse lote	Respondents who a	nswered SMC6_2=1					(100)
ontent			<u>Code</u>		Sample	Population	SMC6_3<=996
'EARS			5 - 85		17,248	5,718,347	
IOT APPLICABLE			996		64,077	22,778,375	SMC6_31
ON'T KNOW			997		194	56,590	
EFUSAL			998		33	19,581	
IOT STATED			999		252	68,842	
		Total			81,804	28,641,735	
/arlable Name	SMC6_4	Length	3	Position	933 •	935	(1)
Question Name	SMK-Q4						
Soncept	Number of cigarette	99					
Question	How many cigarette						
Jniverse	Respondents who a						
lote							SMC6_4<=996
<u>Content</u>			<u>Code</u>		<u>Sample</u>	Population	_
NUMBER OF CIGA	RETTES		1 - 99		17,354	5,767,704	SMC6_41
NOT APPLICABLE			996		64,077	22,778,375	
DON'T KNOW			997		88	21,936	
REFUSAL			998		33	4,878	
NOT STATED			999		252	68,842	
		Total			81,804	28,641,735	
/ariable Name	WT66	Leng	th	8.2	Position	1228 - 1235	
Question Name							Weight used in
Concept	Sampling weig	nt for selected respon	dent				all analysis
Question							
Jniverse	All respondents						
Note	See document	ation on weighting.					

APPENDIX B

Results of contingency table analysis.

Variable Name & Meaning	χ^2 test	d.f.	Statistic value	P value
PRC6_CUR (Province of residence)	Pearson Chi-Square LLR Chi-Square	9 9	22.3672 23.4992	0.0078 0.0052
GE36GURB (Derived rural and urban area – grouped)	Pearson Chi-Square LLR Chi-Square	2 2	21.2294 22.2515	<.0001 <.0001
GE36GCMA (Derived 1991 Census Metropolitan Area – grouped)	Pearson Chi-Square LLR Chi-Square	2 2	4.1918 4.1365	0.1230 0.1264
GE36GHLR (Derived health areas – 26 groups – grouped)	Pearson Chi-Square LLR Chi-Square	26 26	103.1537 99.6342	<.0001 <.0001
GE36GHRO (Derived health areas – 33 groups – grouped)	Pearson Chi-Square LLR Chi-Square	33 33	122.2269 116.5697	<.0001 <.0001
DHC6GAGE (Age – grouped)	Pearson Chi-Square LLR Chi-Square	18 18	1012.3583 943.1849	<.0001 <.0001
DHC6_SEX (Gender)	Pearson Chi-Square LLR Chi-Square	1 1	50.2127 50.2966	<.0001 <.0001
DHC6GHSZ (Derived household size – grouped)	Pearson Chi-Square LLR Chi-Square	4 4	5.9336 5.8921	0.2042 0.2074
DHC6GLE5 (Derived persons <= 5 years old in household – grouped)	Pearson Chi-Square LLR Chi-Square	1 1	0.3673 0.3662	0.5445 0.5451
DHC6G611 (Derived persons 6 to 11 years old in hhld – grouped)	Pearson Chi-Square LLR Chi-Square	1 1	36.6438 35.8356	<.0001 <.0001
HWC6G3KG (Height – adults and children – grouped)	Pearson Chi-Square LLR Chi-Square	13 13	394.9805 380.9162	<.0001 <.0001
DHC6DLVG (Derived living arrangements of the selected respondent)	Pearson Chi-Square LLR Chi-Square	9 9	610.7598 608.6087	<.0001 <.0001
HWC6GSW (Weight in kilograms - grouped)	Pearson Chi-Square LLR Chi-Square	44	398.7164 393.4118	<.0001 <.0001
HWC6GBMI (Derived Body Mass Index (1 decimal place) - grouped)	Pearson Chi-Square LLR Chi-Square	5 5	447.3341 439.5541	<.0001 <.0001

LFC6GO21 (Occupation Codes for main job	Pearson Chi-Square	21	395.7104	<.0001
- 21 groups – grouped)	LLR Chi-Square	21	425.7088	<.0001
LFC6GI13	Pearson Chi-Square	13	312.7330	<.0001
(Industry Codes for main job - 13 groups – grouped)	LLR Chi-Square	13	335.4328	<.0001
PAC6DEE (Derived energy expenditure (1	Pearson Chi-Square	4	222.8556	<.0001
(Derived energy expenditure (1 decimal point))	LLR Chi-Square	4	203.3757	<.0001
PAC6DLEI	Pearson Chi-Square	2	228.8690	<.0001
(Derived participant in leisure physical activity)	LLR Chi-Square	2	209.4993	<.0001
PAC6DFM			200.04((. 0001
(Derived monthly freq. of physical activity lasting > 15	Pearson Chi-Square LLR Chi-Square	6 6	300.9466 286.6957	<.0001 <.0001
min.)				
PAC6DFR	Pearson Chi-Square	3	254.7385	<.0001
(Derived frequency of all physical activity)	LLR Chi-Square	3	237.6286	<.0001
PAC6DFD	Pearson Chi-Square	2	292.1264	<.0001
(Derived participation / daily phys. activities > 15 min.)	LLR Chi-Square	2	275.1194	<.0001
PAC6DPAI	Pearson Chi-Square	3	306.5320	<.0001
(Derived physical activity index)	LLR Chi-Square	3	290.5368	<.0001
SDC6GCB	Pearson Chi-Square	3	387.1970	<.0001
(Country of birth - grouped)	LLR Chi-Square	3	498.4626	<.0001
SDC6GRAC	Pearson Chi-Square	1	38.9072	<.0001
(Derived race or colour – grouped)	LLR Chi-Square	1	41.4480	<.0001
SMC6_4	Pearson Chi-Square	54	97.8939	0.0002
(Number of cigarettes smoked each day - daily smoker)	LLR Chi-Square	54	94.8793	0.0005
CCC6_1H	Pearson Chi-Square	1	2316.2080	<.0001
(Has chronic bronchitis or emphysema)	LLR Chi-Square	1	1391.1395	<.0001
DHC6GBED	Pearson Chi-Square	5	41.8390	<.0001
(Derived number of bedrooms in dwelling – grouped)	LLR Chi-Square	5	40.9440	<.0001
DHC6GECF	Beerson Chi Sauga	0	225 6656	< 0001
(Derived household type –	Pearson Chi-Square LLR Chi-Square	8	235.6656 231.6044	<.0001 <.0001
grouped) HC6GMAR				
(Marital status – grouped)	Pearson Chi-Square LLR Chi-Square	$\begin{vmatrix} 2\\ 2 \end{vmatrix}$	688.2057 687.5607	<.0001 <.0001
HWC6GHT	· · · · · ·			
(Derived standard weight –	Pearson Chi-Square LLR Chi-Square	6 6	383.8739 355.0966	<.0001 <.0001
grouped)		0	555.0900	<.0001

Appendix C

Program used to extract data of NPHS from H356.txt in abbreviated form.

proc format;	
VALUE ACCQ71FM	
1 = ' < THAN 1 YR AGO'	
2 = '1 TO < 2 YEARS AGO'	
3 = '2 - <3 YEARS AGO'	
4 = '3 OR + YEARS AGO'	
5 = 'NEVER'	
6 = 'NOT APPLICABLE'	
7 = 'DON''T KNOW'	
8 = 'REFUSAL'	
9 = 'NOT STATED'	
; VALUE ACCQ72FM	
1 = ' > THAN ONCE A YR'	
2 = 'ONCE A YEAR'	
3 = 'EVERY 2 YEARS'	
4 = 'EVERY 3 YEARS'	
5 = ' < THAN EVERY 3 YR'	
6 = 'NOT APPLICABLE'	
7 = 'DON''T KNOW'	
8 = 'REFUSAL'	
9 = 'NOT STATED'	
;	
;	
;	
, VALUE YEARFM	
9996 = 'NOT APPLICABLE'	
9997 = 'DON''T KNOW'	
9998 = 'REFUSAL'	
9999 = 'NOT STATED'	
;	
, VALUE YESNOFM	
1 = 'YES'	
2 = 'NO'	
6 = 'NOT APPLICABLE'	
7 = 'DON''T KNOW'	
8 = 'REFUSAL'	
9 = 'NOT STATED'	
;	
run;	
<pre>data 'd:\public\h962';</pre>	
<pre>INFILE 'E:\SAS\H356.txt' LRECL = 1243;</pre>	
INPUT	
@ 1 AM66_RNO 6.	
@ 7 PRC6_CUR 2.	
@ 9 GE36GURB 1.	
@ 10 GE36GCMA 3.	

@ 13 @ 17 @ 21 @ 22 @ 24 @ 26	GE36GHLR GE36GHRO DHC6_OWN DHC6GBED INC6G2 INC6DIA5 :	4. 4. 1. 2. 2. 1.
@1222 @1223 @1224 @1225 @1226 @1227 @1228 @1236 ;	VSP6_7 VSP6_8 VSP6_9 VSP6_10 COP6_1 COP6_2 WT66 WT66_N	1. 1. 1. 1. 1. 8.2 8.2
label AM66_RNC PRC6_CUR GE36GURE GE36GCMA : :	= 'Province = 'Derived r	mber on Health Microdata file' of residence' rural and urban area – grouped' 991 Census Metropolitan Area – grouped'
VSP6_7 VSP6_8 WT66	= 'Number of	times attacked at school / school bus' times verbally abused outside of school' weight for selected respondent'
format PRC6_CUF GE36GURE GE36GCMA GE36GHLF : : RSS6_9 RSS6_10	PROVFM. DVGURBFM. DVGCMAFM. DVGHLRFM. YESNOFM.	
RSS6_10 VSP6_1 VSP6_2 VSP6_4 VSP6_5 COP6_1 COP6_2	RSS_FREQ. OFTENFM. OFTENFM. OFTENFM. VSP_NUM. GOOD5FM. GOOD5FM. ;	

run;

APPENDIX D

Programs used for contingency table analysis.

```
Program 1.
data 'e:\temp4\data_a';
Set 'd:\public\h962';
keep wt66;
if ccc6_1c<=2;
run;
proc means data='e:\temp4\data_a' noprint;
var WT66;
output out='e:\temp4\aaa' mean=mwt66;
run;
proc print;
run;
Program 2.
data 'e:\temp4\data';
Set 'd:\public\h962';
Keep
PRC6_CUR
                   PAC6DFM
GE36GURB
                   PAC6DFR
GE36GCMA
                  PAC6DFD
GE36GHLR
                  PAC6DPAI
GE36GHRO
                  DGC6_1G
DHC6_OWN
                  DGK6_1
DHC6GBED
                   SMC6_3
DHC6GAGE
                   SMC6_4;
DHC6_SEX
DHC6GMAR
DHC6GHSZ
DHC6GLE5
DHC6G611
DHC6GECF
DHC6DLVG
CCC6_1C
CCC6_C5
CCC6_C6
CCC6_1H
SDC6GCB
SDC6GRAC
LFC6G021
LFC6GI13
HWC6GHT
HWC6G3KG
HWC6GBMI
HWC6GSW
HWS6_1
HWS6_5
PAC6DEE
PAC6DLEI
```

```
Run;
data 'e:\temp4\data';
set 'e:\temp4\data';
if CCC6_1C<=2;
Wt66=Wt66/350.21055985;
run;
Program 3.
proc freq data='e:\temp4\data';
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables(PRC6_CUR
       GE36GURB
       GE36GCMA
       GE36GHLR
       GE36GHRO
       DHC6GAGE
       DHC6 SEX
       DHC6GHSZ
       DHC6GLE5
       DHC6G611) *CCC6_1C/chisg ;
*output out='e:\temp1\tfreq0';
run;
options nonumber nodate pagesize=5000;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 HWC6G3KG;
if HWC6G3KG<=140;
if 0<=HWC6G3KG<=10 then HWC6G3KG=1;
if 10<HWC6G3KG<=20 then HWC6G3KG=2;
if 20<HWC6G3KG<=30 then HWC6G3KG=3;
if 30<HWC6G3KG<=40 then HWC6G3KG=4;
if 40<HWC6G3KG<=50 then HWC6G3KG=5;
if 50<HWC6G3KG<=60 then HWC6G3KG=6;
if 60<HWC6G3KG<=70 then HWC6G3KG=7;
if 70<HWC6G3KG<=80 then HWC6G3KG=8;
if 80<HWC6G3KG<=90 then HWC6G3KG=9;
if 90<HWC6G3KG<=100 then HWC6G3KG=10;
if 100<HWC6G3KG<=110 then HWC6G3KG=11;
if 110<HWC6G3KG<=120 then HWC6G3KG=12;
if 120<HWC6G3KG<=130 then HWC6G3KG=13;
if 130<=HWC6G3KG<=140 then HWC6G3KG=14;
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables HWC6G3KG*CCC6_1C/Chisq;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 HWC6GBMI ;
if HWC6GBMI <=99.6;
if 10<HWC6GBMI<=20 then HWC6GBMI=1;
if 20<HWC6GBMI<=30 then HWC6GBMI=2;
```

```
if 30<HWC6GBMI<=40 then HWC6GBMI=3;
if 40<HWC6GBMI<=50 then HWC6GBMI=4;
if 50<HWC6GBMI<=60 then HWC6GBMI=5;
run:
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables HWC6GBMI*CCC6_1C/Chisg;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66
                  HWC6GSW ;
if
    HWC6GSW <=6;
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables HWC6GSW*CCC6_1C/Chisq;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 HWC6GHT ;
if
     HWC6GHT <=64;
if 0<=HWC6GHT<=10 then HWC6GHT=1;
if 10<HWC6GHT<=20 then HWC6GHT=2;
if 20<HWC6GHT<=30 then HWC6GHT=3;
if 30<HWC6GHT<=40 then HWC6GHT=4;
if 40<HWC6GHT<=50 then HWC6GHT=5;
if 50<HWC6GHT<=60 then HWC6GHT=6;
if 60<HWC6GHT<=70 then HWC6GHT=7;
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables HWC6GHT*CCC6_1C/Chisq;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 LFC6GO21 ;
if LFC6G021 <=96;
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables LFC6G021*CCC6_1C/Chisq;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 LFC6GI13 ;
if
    LFC6GI13 <=96;
run;
proc freq data=freq ;
weight WT66;
```

```
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables LFC6GI13*CCC6 1C/Chisg;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 PAC6DEE ;
     PAC6DEE <=99.6;
if
if 0<=PAC6DEE<=10 then PAC6DEE=1;
if 10<PAC6DEE<=20 then PAC6DEE=2;
if 20<PAC6DEE<=30 then PAC6DEE=3;
if 30<PAC6DEE<=40 then PAC6DEE=4;
run:
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables PAC6DEE*CCC6_1C/Chisq;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 PAC6DLEI ;
if
      PAC6DLEI <=6;
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables PAC6DLEI*CCC6_1C/Chisq;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 PAC6DFM ;
if PAC6DFM <=996;
if 0<=PAC6DFM<=5 then PAC6DFM=1:
if 5<PAC6DFM<=10 then PAC6DFM=2;
if 10<PAC6DFM<=15 then PAC6DFM=3;
if 15<PAC6DFM<=20 then PAC6DFM=4;
if 20<PAC6DFM<=25 then PAC6DFM=5;
if 25<PAC6DFM<=255 then PAC6DFM=6;
if 255<PAC6DFM then PAC6DFM=7;
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables PAC6DFM*CCC6_1C/Chisq;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 PAC6DFR ;
if
    PAC6DFR \leq =6;
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables PAC6DFR*CCC6_1C/Chisq;
```

data freq; set 'e:\temp4\data'; keep CCC6_1C WT66 PAC6DFD; if PAC6DFD<=6; run; proc freq data=freq ; weight WT66; title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA"; tables PAC6DFD*CCC6_1C/Chisq; run; data freq; set 'e:\temp4\data'; keep CCC6_1C WT66 PAC6DPAI; if PAC6DPAI<=6; run; proc freq data=freq ; weight WT66; title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA"; tables PAC6DPAI*CCC6_1C/Chisq; run: data freq; set 'e:\temp4\data'; keep CCC6_1C WT66 SDC6GCB; if SDC6GCB<=4; run; proc freq data=freq ; weight WT66; title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA"; tables SDC6GCB*CCC6_1C/Chisq; run: data freq; set 'e:\temp4\data'; keep CCC6_1C WT66 SDC6GRAC; if SDC6GRAC<=2; run; proc freq data=freq ; weight WT66; title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA"; tables SDC6GRAC*CCC6_1C/Chisq; run; data freq; set 'e:\temp4\data'; keep CCC6_1C WT66 SMC6_3; if SMC6_3<=996; if 0<=SMC6_3<=10 then SMC6 3=1; if 10<SMC6_3<=20 then SMC6_3=2; if 20<SMC6_3<=30 then SMC6_3=3; if 30<SMC6_3<=40 then SMC6_3=4; if 40<SMC6_3<=50 then SMC6_3=5; if 50<SMC6_3<=60 then SMC6_3=6; if 60<SMC6_3<=70 then SMC6_3=7;

run;

```
if 60<SMC6_3<=70 then SMC6_3=7;
   else SMC6_3=8;
run;
proc freq data=freq ;
weight WT66;
title "FREOUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables SMC6_3*CCC6_1C/Chisq;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 SMC6_4;
if SMC6_4<=996;
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables SMC6_4*CCC6_1C/Chisq;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 CCC6_1H;
if CCC6_1H<=2;
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables CCC6_1H*CCC6_1C/Chisq;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 DHC6GBED;
if 0<=DHC6GBED<=5;
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables DHC6GBED*CCC6_1C/Chisq;
run;
data freg;
set 'e:\temp4\data';
keep CCC6_1C WT66 DHC6GMAR;
if 0<=DHC6GMAR<=3;
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables DHC6GMAR*CCC6_1C/Chisq;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 DHC6GECF;
if 0<=DHC6GECF<=9;
```

```
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables DHC6GECF*CCC6_1C/Chisq;
run;
data freq;
set 'e:\temp4\data';
keep CCC6_1C WT66 DHC6DLVG;
if 0<=DHC6DLVG<=10;
run;
proc freq data=freq ;
weight WT66;
title "FREQUENCY TABLE ABOUT VARIABLES RELATED TO ASTHMA";
tables DHC6DLVG*CCC6_1C/Chisq;
run;
```

APPENDIX E

Programs for trend test.

```
data trend;
SET 'e:\temp4\data';
KEEP CCC6_1C WT66 DHC6GAGE ;
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*DHC6GAGE/trend NOPRINT ;
RUN;
/******
data trend;
SET 'e:\temp4\data';
KEEP CCC6_1C WT66 DHC6GBED ;
IF 0<=DHC6GBED<=99;
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*DHC6GBED/trend NOPRINT ;
RUN;
/**************
data trend;
SET 'e:\temp4\data';
KEEP CCC6_1C WT66 DHC6GHSZ ;
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*DHC6GHSZ/trend NOPRINT ;
RUN;
/**************
data trend;
SET 'e:\temp4\data';
KEEP PAC6DEE CCC6_1C WT66;
IF PAC6DEE<=34.4;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*PAC6DEE/trend noprint;
RUN;
/**************
data trend;
SET 'e:\temp4\data';
KEEP PAC6DFM CCC6_1C WT66;
IF PAC6DFM<=255;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*PAC6DFM/trend NOPRINT ;
RUN;
/******
data trend;
```

SET 'e:\temp4\data';

```
KEEP PAC6DFR CCC6_1C WT66;
IF PAC6DFR<=3;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*PAC6DFR/trend NOPRINT ;
RUN;
/******
data trend;
SET 'e:\temp4\data';
KEEP PAC6DPAI CCC6_1C WT66 SMC6_3 ;
IF SMC6_3<=85;
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*SMC6_3/TREND NOPRINT ;
RUN;
/*************/
data trend;
SET 'e:\temp4\data';
KEEP CCC6_1C WT66 SMC6_4 ;
IF 0<=SMC6_4<=99;
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*SMC6_4/trend NOPRINT ;
RUN;
/*************/
data trend;
SET 'e:\temp4\data';
KEEP CCC6_1C WT66 SMC6_4 ;
IF 0<=SMC6_4<=15;
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*SMC6_4/trend NOPRINT ;
RUN;
/**************/
data trend;
SET 'e:\temp4\data';
KEEP CCC6_1C WT66 SMC6_4 ;
IF 15<=SMC6_4<=98;
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*SMC6_4/trend NOPRINT ;
RUN;
/*************
data trend;
SET 'e:\temp4\data';
KEEP CCC6_1C WT66 HWC6GHT ;
IF 0<=HWC6GHT<99;
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*HWC6GHT/trend NOPRINT ;
RUN;
```

```
/**************/
data trend;
SET 'e:\temp4\data';
KEEP CCC6_1C WT66 HWC6GSW ;
IF 0<=HWC6GSW<6;</pre>
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*HWC6GSW/trend NOPRINT ;
RUN;
/*****
data trend;
SET 'e:\temp4\data';
KEEP CCC6_1C WT66 HWC6GSW ;
IF 0<=HWC6GSW<=3;</pre>
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*HWC6GSW/trend NOPRINT ;
RUN;
/**************/
data trend;
SET 'e:\temp4\data';
KEEP CCC6_1C WT66 HWC6G3KG ;
IF 0<=HWC6G3KG<=137;
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*HWC6G3KG/trend NOPRINT ;
RUN;
/****/
data trend;
SET 'e:\temp4\data';
KEEP CCC6_1C WT66 HWC6G3KG ;
IF 70<=HWC6G3KG<=137;
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*HWC6G3KG/trend NOPRINT ;
RUN;
/**********
data trend;
SET 'e:\temp4\data';
KEEP CCC6_1C WT66 HWC6GBMI ;
IF 0<=HWC6GBMI<=60;</pre>
RUN;
PROC FREQ DATA=TREND;
WEIGHT WT66;
TABLES CCC6_1C*HWC6GBMI/trend NOPRINT ;
RUN;
/*****/
```

APPENDIX F

Program used for odds ratio and relative risk analysis.

```
data data;
set 'e:\temp4\data';
keep CCC6_1C WT66 PRC6_CUR ;
RUN;
/*****
DATA DATA;
SET DATA;
if PRC6_CUR=10 THEN NFLD=0; ELSE NFLD=1;
IF PRC6_CUR=11 THEN PEI=0; ELSE PEI=1;
IF PRC6_CUR=12 THEN NS=0; ELSE NS =1;
IF PRC6_CUR=13 THEN NB=0; ELSE NB=1;
IF PRC6_CUR=24 THEN QUE=0; ELSE QUE=1;
IF PRC6_CUR=35 THEN ONT=0; ELSE ONT=1;
IF PRC6_CUR=46 THEN MB=0; ELSE MB=1;
IF PRC6_CUR=47 THEN SASK=0; ELSE SASK=1;
IF PRC6_CUR=48 THEN AB=0; ELSE AB=1;
IF PRC6_CUR=59 THEN BC=0; ELSE BC=1;
RUN;
PROC FREQ DATA=DATA;
weight WT66;
tables (NFLD PEI NS NB QUE ONT MB SASK AB BC)*CCC6_1C
       /chisq CMH ALPHA=0.01 noprint;
run;
************************
DATA TEST;
SET DATA;
IF PRC6_CUR=12 OR PRC6_CUR=10;
IF PRC6_CUR=10 THEN PRC6_CUR=100;
PROC FREQ DATA=TEST;
WEIGHT WT66;
TABLES PRC6_CUR*CCC6_1C/CHISQ MEASURE ALPHA=0.01;
RUN;
DATA TEST;
SET DATA;
IF PRC6_CUR=12 OR PRC6_CUR=35 OR PRC6_CUR=24 THEN HIGH_LOW=0; ELSE
HIGH_LOW=1;
PROC FREQ DATA=TEST;
WEIGHT WT66;
TABLES HIGH_LOW*CCC6_1C/CHISQ MEASURE ALPHA=0.01;
RUN:
****************************
data data;
set 'e:\temp4\data';
keep CCC6_1C WT66 GE36GURB ;
IF GE36GURB<=2;
```

RUN; PROC FREQ DATA=DATA; weight WT66; TABLES GE36GURB*CCC6_1C/CHISQ MEASURES ALPHA=0.01; RUN: ***************************** data data; set 'e:\temp4\data'; keep CCC6_1C WT66 LFC6GI13 ; IF LFC6GI13<=13;</pre> RUN; DATA TEST; SET DATA; IF LFC6GI13=1 OR LFC6GI13=11; IF LFC6GI13=11 THEN LFC6GI13=0; PROC FREQ DATA=TEST; WEIGHT WT66; TABLES LFC6GI13*CCC6_1C/CHISQ MEASURE ALPHA=0.01; RUN; DATA TEST; SET DATA; IF LFC6GI13=9 OR LFC6GI13=10 OR LFC6GI13=11 THEN HIGH_LOW=0; ELSE HIGH_LOW=1; PROC FREQ DATA=TEST; WEIGHT WT66; TABLES HIGH_LOW*CCC6_1C/CHISQ MEASURE ALPHA=0.01; RUN; ****** data data; set 'e:\temp4\data'; keep CCC6_1C WT66 PAC6DEE PAC6DLEI PAC6DFR PAC6DFD PAC6DPAI; IF PAC6DEE<=99.6; RUN; /***** DATA TEST; SET DATA; IF LFC6GI13=1 OR LFC6GI13=11; IF LFC6GI13=11 THEN LFC6GI13=0; PROC FREQ DATA=TEST; WEIGHT WT66; TABLES LFC6GI13*CCC6_1C/CHISQ MEASURE ALPHA=0.01; RUN; *****/ DATA TEST; SET DATA; IF PAC6DEE=99.6 THEN CH AD=0; ELSE CH AD=1; PROC FREQ DATA=TEST; WEIGHT WT66; TABLES CH_AD*CCC6_1C/CHISQ MEASURE ALPHA=0.01; RUN; DATA TEST; SET DATA;

IF PAC6DFR<=3; IF PAC6DFR>1 THEN PAC6DFR=4; PROC FREQ DATA=TEST; WEIGHT WT66; TABLES PAC6DFR*CCC6_1C/CHISQ MEASURE ALPHA=0.01; RUN; DATA TEST; SET DATA; IF PAC6DPAI<=3; IF PAC6DPAI>1 THEN PAC6DPAI=4; PROC FREQ DATA=TEST; WEIGHT WT66; TABLES PAC6DPAI*CCC6_1C/CHISQ MEASURES ALPHA=0.01; RUN: DATA TEST; SET DATA; IF PAC6DFD<=2; PROC FREQ DATA=TEST; WEIGHT WT66: TABLES PAC6DFD*CCC6_1C/CHISQ MEASURE ALPHA=0.01; RUN; *********************** data data; set 'e:\temp4\data'; keep CCC6_1C WT66 SDC6GCB SDC6GRAC DHC6GMAR; RUN; DATA TEST; SET DATA; SDC6GCB= 1 THEN SDC6G1 IF =0; ELSE SDC6G1 =1; THEN SDC6G2 ELSE SDC6G2 IF SDC6GCB= 2 =0; =1;THEN SDC6G3 IF SDC6GCB= 3 =0; ELSE SDC6G3 =1;SDC6GCB= THEN SDC6G4 =0; ELSE SDC6G4 Δ =1; IF PROC FREO DATA=TEST; WEIGHT WT66; TABLES (SDC6G1-SDC6G4) *CCC6_1C/CHISQ MEASURE ALPHA=0.01; RUN; DATA TEST; SET DATA; if SDC6GRAC<=2; PROC FREQ DATA=TEST; WEIGHT WT66; TABLES (SDC6grac) *CCC6_1C/CHISQ MEASURE ALPHA=0.01; RUN; DATA TEST; SET DATA; if DHC6GMAR<=3; if DHC6GMAR ne 2 then DHC6GMAR=4; PROC FREQ DATA=TEST; WEIGHT WT66; TABLES (DHC6GMAR) *CCC6_1C/CHISQ MEASURE ALPHA=0.01; RUN; DATA TEST;

SET DATA; if DHC6GMAR=1 or DHC6GMAR=3; if DHC6GMAR=1 then DHC6GMAR=4; PROC FREQ DATA=TEST; WEIGHT WT66; TABLES (DHC6GMAR) *CCC6_1C/CHISQ MEASURE ALPHA=0.01; RUN; **************************** data data; set 'e:\temp4\data'; keep CCC6_1C WT66 smc6_4 smc6_3 dhc6gage; if smc6_3<=996; RUN; data test; set data; if smc6_3<996 then smoke=1; else smoke=2; proc freq data=test; weight wt66; tables smoke*ccc6_1c/chisq measures alpha=0.01; run; data test; set data; if smc6 3 ne 996; if smc6_3<20 then smoke=1; else smoke=2; proc freq data=test; weight wt66; tables smoke*ccc6_1c/chisq measures alpha=0.01; run; data test; set data; if smc6_4 ne 996; if smc6_4<10 then smoke=1; else smoke=2; proc freq data=test; weight wt66; tables smoke*ccc6_1c/chisq measures alpha=0.01; run; data test; set data; if smc6_4 ne 996 and dhc6gage<=6; if smc6_4>10 then smoke=1; else smoke=2; proc freq data=test; weight wt66; tables smoke*ccc6_1c/chisq measures alpha=0.01; run; data test; set data; if smc6_4 ne 996 and dhc6gage>6; if smc6_4>10 then smoke=1; else smoke=2; proc freq data=test; weight wt66; tables smoke*ccc6_1c/chisq measures alpha=0.01; run;

```
*****************************
data data;
set 'e:\temp4\data';
keep CCC6_1C WT66 dhc6_own DHC6GBED;
RUN;
data test;
set data;
if dhc6_own<=2 ;
IF dhc6_own=1 THEN dhc6_own=3;
proc freq data=test;
weight wt66;
tables dhc6_own*ccc6_1c/chisq measures alpha=0.01;
run;
************************
data data;
set 'e:\temp4\data';
keep CCC6_1C WT66 DHC6_SEX CCC6_1H HWC6GSW DHC6GAGE ;
RUN;
data test;
set data;
*if dhc6_own<=2 ;</pre>
IF dhc6_SEX=1 THEN dhc=4;ELSE DHC=3;
proc freq data=TEST;
weight wt66;
tables DHC*ccc6_1c/chisq measures alpha=0.01;
run;
data test;
set data;
if dhc6GAGE<=4 ;
*IF dhc6_SEX=1 THEN dhc=4;*ELSE DHC=3;
proc freq data=TEST;
weight wt66;
tables DHC6_SEX*ccc6_1c/chisq measures alpha=0.01;
run;
data test;
set data;
if dhc6GAGE>4 ;
IF dhc6_SEX=1 THEN dhc=4; ELSE DHC=3;
proc freq data=TEST;
weight wt66;
tables DHC*ccc6_1c/chisq measures alpha=0.01;
run;
data test;
set data;
if HWC6GSW<=4 ;
IF HWC6GSW<=3 THEN DHC=1;ELSE DHC=0;
proc freq data=TEST;
weight wt66;
tables DHC*ccc6_1c/chisq measures alpha=0.01;
run;
```

APPENDIX G

Programs used to fit loglinear regression model.

```
data loglin;
set 'e:\temp4\data' ;
if 0<=PRC6_CUR<=59
and GE36GURB<=6
AND HWC6G3KG<=138
AND HWC6GBMI<=99.6
AND HWC6GSW<=6
AND HWC6GHT<=64
AND HWS6_1<=6
AND HWS6_5<=6
AND LFC6G021<=96
AND LFC6GI13<=96
AND PAC6DEE<=99.6
AND PAC6DLEI<=6
AND PAC6DFM<=996
AND PAC6DFR<=6
AND PAC6DFD<=6
AND PAC6DPAI<=9
AND SDC6GCB<=4
AND SDC6GRAC<=2
AND SMC6_3<=996
AND SMC6_4<=996
AND DGC6_1G<=6
AND CCC6_1H<=2
AND DGK6_1<=6
AND DHC6_OWN<=2
AND DHC6GBED<=5
AND DHC6GMAR<=3
AND DHC6GECF<=99
AND DHC6DLVG<=10;
run;
data new;
set loglin;
*keep CCC6_1C WT66 PRC6_CUR LFC6GI13 PAC6DPAI
      SMC6_3 DHC6_OWN DHC6_SEX DHC6GAGE;
IF PRC6_CUR=12 OR PRC6_CUR=24 OR PRC6_CUR=35
   THEN PRC6_CUR=0; ELSE PRC6_CUR=1;
*IF 9<=LFC6GI13<=11 THEN LFC6GI13=0;* ELSE LFC6GI13=1;
IF 9<=LFC6GI13<=11 OR LFC6GI13=6 OR LFC6GI13=8 OR 12<=LFC6GI13<=13
   THEN LFC6GI13=1;
     ELSE IF LFC6GI13=96 THEN LFC6GI13=2;
       ELSE LFC6GI13=0;
IF PAC6DPAI=1 THEN PAC6DPAI=0;
   ELSE IF PAC6DPAI>1 AND PAC6DPAI NE 6 THEN PAC6DPAI=1;
      ELSE PAC6DPAI=2;
IF SMC6_3=996 THEN SMC6_3=0; ELSE SMC6_3=1;
IF SDC6GCB>1 THEN SDC6GCB=0; ELSE SDC6GCB=1;
```

IF DHC6GBED<=2 THEN DHC6GBED=1; ELSE DHC6GBED=0; RUN; PROC CATMOD DATA=NEW; WEIGHT WT66; MODEL CCC6_1C*PRC6_CUR* LFC6GI13 *PAC6DPAI *DHC6_SEX *SMC6_3 *DHC6_OWN =_RESPONSE_ /NOPROFILE NORESPONSE NOITER NOPARM; LOGLIN CCC6_1C|PRC6_CUR| LFC6GI13 |PAC6DPAI |DHC6_SEX |SMC6_3 |DHC6_OWN @5; RUN;

APPENDIX H

Programs used to fit logistic regression model for all respondents.

data logistic; set 'e:\temp4\data' ; if 0<=PRC6 CUR<=59 and GE36GURB<=6 AND HWC6G3KG<=138 AND HWC6GBMI<=99.6 AND HWC6GSW<=6 AND HWC6GHT<=64 AND HWS6_1<=6 AND HWS6_5<=6 AND LFC6G021<=96 AND LFC6GI13<=96 AND PAC6DEE<=99.6 AND PAC6DLEI<=6 AND PAC6DFM<=996 AND PAC6DFR<=6 AND PAC6DFD<=6 AND PAC6DPAI<=9 AND SDC6GCB<=4 AND SDC6GRAC<=2 AND SMC6_3<=996 AND SMC6_4<=996 AND DGC6 1G<=6 AND CCC6_1H<=2 AND DGK6_1<=6 AND DHC6 OWN<=2 AND DHC6GBED<=5 AND DHC6GMAR<=3 AND DHC6GECF<=99 AND DHC6DLVG<=10; *if PRC6_CUR=10 THEN NFLD=0; *ELSE NFLD=0; IF PRC6 CUR=11 THEN PEI=1; ELSE PEI=0; IF PRC6_CUR=12 THEN NS=1; ELSE NS =0; IF PRC6_CUR=13 THEN NB=1; ELSE NB=0; IF PRC6_CUR=24 THEN QUE=1; ELSE QUE=0; IF PRC6_CUR=35 THEN ONT=1; ELSE ONT=0; IF PRC6_CUR=46 THEN MB=1; ELSE MB=0; IF PRC6_CUR=47 THEN SASK=1; ELSE SASK=0; IF PRC6_CUR=48 THEN AB=1; ELSE AB=0; IF PRC6_CUR=59 THEN BC=1; ELSE BC=0; IF 0<=HWC6GBMI<=57.9 THEN HWC6GB1 =HWC6GBMI; ELSE HWC6GB1=0; IF HWC6GSW=1 THEN HWC6G1 =1; ELSE HWC6G1 =0;IF HWC6GSW=2 THEN HWC6G2 =1; ELSE HWC6G2 =0;IF HWC6GSW=3 THEN HWC6G3 ELSE HWC6G3 =0; =1; IF HWC6GSW=4 THEN HWC6G4 =1; ELSE HWC6G4 =0;HWS6_1=1 THEN HWS1 =1; ELSE HWS1 =0; HWS6_1=2 THEN HWS2 =1; ELSE HWS2 =0; ΤF IF

IF	HWS6_1=3	THEN	HWS3	=1;	ELSE	HWS3	=0;			
* IF	0<=HWS6_2KG HWS6KG1	<=185 =0;	THEN	HWS6K	G1	=HWS6	_2KG;	* ELSI	3	
IF	HWS6_5=	1	THEN	HWS61	=1;	ELSE	HWS61	=0:		
IF	HWS6_5=	2	THEN		=1;		HWS62			
**	111100_0	-	11121		- 1			• /		
IF	LFC6GI13=	1	THEN	LFC1		ELSE	LFC1	=0;		
IF	LFC6GI13=	2	THEN	LFC2	=1;	ELSE	LFC2	=0;		
IF	LFC6GI13=	3	THEN	LFC3		ELSE	LFC3	=0;		
IF	LFC6GI13=	4	THEN	LFC4		ELSE	LFC4	=0;		
IF	LFC6GI13=	5	THEN	LFC5		ELSE	LFC5	=0;		
IF	LFC6GI13=	6	THEN	LFC6		ELSE		=0;		
IF	LFC6GI13=	7	THEN	LFC7		ELSE		=0;		
IF	LFC6GI13=	8	THEN	LFC8		ELSE		=0;		
IF	LFC6GI13=	9 10	THEN THEN	LFC9 LFC10		ELSE	LFC9 LFC10			
IF IF	LFC6GI13= LFC6GI13=	11	THEN		-⊥; =1;	ELSE ELSE	LFC10 LFC11			
IF	LFC6GI13=	12	THEN		=1;	ELSE	LFC12			
IF	LFC6GI13=	13	THEN	LEC13	-1;	ELSE	LFC13			
ΤΓ	DLCOGII2-	10	IUCN	лгстэ	- 1 ;	2022	пстр	-0;		
IF	PAC6DLEI=1				=1;		PAC6D		=0;	
IF	PAC6DLEI=2	THEN	PAC6D	L2	=1;	ELSE	PAC6D	L2	=0;	
IF	0 <pac6dfm<= ELSE if</pac6dfm<= 		THEN M=0 th		FM1 6DFM1=			PAC6DFI	M1=0;	
IF	PAC6DPAI=	1	THEN	PAC6D	PA1	=1;	ELSE	PAC6D	PA1	=0;
IF	PAC6DPAI=	2	THEN	PAC6D	PA2	=1;	ELSE	PAC6D	PA2	=0;
IF	PAC6DPAI=	3	THEN	PAC6D	PA3	=1;	ELSE	PAC6D	PA3	=0;
IF	SDC6GCB=	1	THEN	SDC6G		=1;	ELSE	SDC6G		=0;
IF	SDC6GCB=	2	THEN	SDC6G		=1;	ELSE	SDC6G		=0;
IF	SDC6GCB=	3	THEN	SDC6G		=1;	ELSE	SDC6G		=0;
IF	SDC6GCB=	4	THEN	SDC6G	4	=0;	ELSE	SDC6G	4	=0;
IF	SDC6GRAC=	1	THEN	SDC6G	R1	=1;	ELSE	SDC6G	R1	=0;
*IF	SDC6GRAC=	2	THEN	SDC6G	R2	=0;	*ELSE		SDC 60	GR2=0;
IF	SMC6_3=996				31		ELSE		31	=100;
IF	0<=SMC6_3<=	=85 THE	EN	SMC6_	_32	=SMC6	_3;	ELSE	SMC6_	_32=0;
IF	0<=SMC6_4<=	=99 THE	EN	SMC6_	41=SMC	26_4;	ELSE	SMC6_4	1	=0;
*IF	DHC6_OWN=	1	THEN	DHC6_	01	=0;	*ELSE		DHC 6	01=0;
IF	DHC6_OWN=	2	THEN	DHC6_		=1;		DHC6_		_
*IF	DHC6GBED=	0	THEN	DHC6G	ומי	=1;	*ELSE		חשמהנ	GB1=0;
*1F	DHC6GBED=	0 1	THEN				*ELSE			GB1=0; GB2=0;
*IF	DHC6GBED=	2	THEN				*ELSE			GB3=0;
*IF	DHC6GBED=	3	THEN	DHC60			*ELSE			GB4=0;
*IF	DHC6GBED=	4	THEN	DHC60			*ELSE			GB5=0;
*IF	DHC6GBED=	5	THEN	DHC60		-	*ELSE			GB6=0;
IF	0<=DHC6GBEI		THEN	DHC60			GBED;			GB1=0;
							-			-

,

IF IF	DHC6_SEX= DHC6_SEX=	1 2	THEN THEN	DHC6_S1 DHC6_S2	=1; =1;	ELSE ELSE	DHC6_S1 DHC6_S2		=0; =0;
							_		
*IF	DHC6GMAR=	1	THEN	DHC6GM1	=1;		DHC6GM1		
IF	DHC6GMAR=	2	THEN	DHC6GM2	=1;	ELSE	DHC6GM2		=0;
IF	DHC6GMAR=	3	THEN	DHC6GM3	=1;	ELSE	DHC6GM3		=0;
*IF	DHC6GHSZ=	1	THEN	DHC6GH1	=1;	*ELSE	Ľ	HC6GH	H1=0;
*IF	DHC6GHSZ=	2	THEN	DHC6GH2	=1;	*ELSE	Γ	HC6GF	H2=0;
*IF	DHC6GHSZ=	3	THEN	DHC6GH3	=1;	*ELSE	E	DHC6GF	I3=0;
*IF	DHC6GHSZ=	4	THEN (DHC6GH4	=1;	*ELSE	E	HC6GF	H4=0;
*IF	DHC6GHSZ=	5	THEN	DHC6GH5	=1;	*ELSE	Γ	OHC 6 GH	H5=0;
*IF	0<=DHC6GHS2	2<=5	THEN	DHC6GH	=1;	*ELSE	Ε	DHC6GH	H=0;
IF	DHC6GLE5=	1	THEN	DHC6GLE1	=1;	ELSE	DHC6GLE	21	=0;
IF	DHC6GLE5=	2	THEN	DHC6GLE2	=1;	ELSE	DHC6GLE		=0;
IF	DHC6G611=	1	THEN	DHC6G61	=1;	ELSE	DHC6G61		=0;
*IF	DHC6G611=	2	THEN	DHC6G62	=1;	*ELSE	DHC6G62	2	=0;
IF	DHC6GECF=	1	THEN	DHC6GE1	=1;	ELSE	DHC6GE1	-	=0;
IF	DHC6GECF=	2	THEN	DHC6GE2	=1;	ELSE	DHC6GE2	2	=0;
IF	DHC6GECF=	3	THEN	DHC6GE3	=1;	ELSE	DHC6GE3	3	=0;
IF	DHC6GECF=	4	THEN	DHC6GE4	=1;	ELSE	DHC6GE4	ł	=0;
IF	DHC6GECF=	5	THEN	DHC6GE5	=1;	ELSE	DHC6GE5	5	=0;
IF	DHC6GECF=	6	THEN	DHC6GE6	=1;	ELSE	DHC6GE6	5	=0;
/*IF	DHC6GECF=	7	THEN	DHC6GE7	=1;	ELSE	DHC6GE7	7	=0;*/
IF	DHC6GECF=	8	THEN	DHC6GE8	=1;	ELSE	DHC6GE8	3	=0;
IF	DHC6GECF=	9	THEN	DHC6GE9	=1;	ELSE	DHC6GE9)	=0;
IF	DHC6DLVG=	1	THEN	DHC6DL1	=1;	ELSE	DHC6DL1	L	=0;
IF	DHC6DLVG=	2	THEN	DHC6DL2	=1;	ELSE	DHC6DL2		=0;
IF	DHC6DLVG=	3	THEN	DHC6DL3	=1;	ELSE	DHC6DL3		=0;
IF	DHC6DLVG=	4	THEN	DHC6DL4	=1;	ELSE	DHC6DL4		=0;
IF	DHC6DLVG=	5	THEN	DHC6DL5	=1;	ELSE	DHC6DL5		=0;
IF	DHC6DLVG=	6	THEN	DHC6DL6	=1;	ELSE	DHC6DL6		=0;
/*IF	DHC6DLVG=	7	THEN	DHC6DL7	=1;	ELSE	DHC6DL7		=0;*/
IF	DHC6DLVG=	8	THEN	DHC6DL8	=1;	ELSE	DHC6DL8		=0;
IF	DHC6DLVG=		THEN		=1;	ELSE	DHC6DL9		=0;
IF	DHC6DLVG=		THEN		=1;		DHC6DL1		=0;
<pre>run;</pre>									·
_	logistic dat	ca =lgt	data I	ESCENDING;					
-	nt WT66;								
	CCC6_1C=								
	PEI NB QUE								
GE36GU1-GE36GU2 HWC6GB1 HWC6G1-HWC6G4 HWS61-HWS62 LFC1-LFC13 PAC6DFM1									
	PA1-PAC6DPA			G3					
	_31 SMC6_32 S			2111					
	O2 DHC6GB1 I								
	M3 DHC6G61 I								
	E8-DHC6GE9								
	DL8-DHC6DL9 I								
	fit SELECTION	JN=STEI	WISE 1	;					
run;									

```
/*
link=logit
link=probit
link=cloglog
proc logistic data =lgtdata DESCENDING;
weight WT66;
model CCC6_1C=
NFLD
     NB
            AB
                  GE36GU1 HWC6G1
HWC6G2
            HWC6G3 LFC1 LFC3 LFC6
LFC9 LFC10 LFC13 SMC6_31 SMC6_41
LFC11 LFC12
                 PAC6DPA1 PAC6DPA2
                SDC6G2
                              SDC6G3
PAC6DPA3 SDC6G1
DHC6 O2
             DHC6_S2
                       DHC6GM2 DHC6GM3
DHC6GE1
             DHC6GE4
                       DHC6GE5 DHC6GE6
DHC6DL5 DHC6DL6 DHC6GAGE
/lackfit link=logit outroc=roc1 roceps=0;
output out=outp p=p;
run;
proc plot data=roc1;
plot (_SENSIT_) *(_1MSPEC_);
run;
```

```
In Splus:
>plot(c(-0.0,1),c(-0.0,1),type='n', xlab='1-Specificity',
ylab='Sensitivity')
>lines(rocl$.1MSPEC,rocl$.SENSIT.)
>lines(c(0,1),c(0,1))
```

APPENDIX I

Programs used for stepwise logistic regression and cross validation

for infants, children and adults.

data lgtdata1; set 'e:\temp4\data'; if PRC6_CUR<=60 and GE36GURB<=6 AND HWC6G3KG<=138 AND HWC6GBMI<=99.6 AND HWC6GSW<=6 AND HWC6GHT<=64 AND HWS6_1<=6 AND HWS6_5<=6 AND LFC6G021<=96 AND LFC6GI13<=96 AND PAC6DEE<=99.6 AND PAC6DLEI<=6 AND PAC6DFM<=996 AND PAC6DFR<=6 AND PAC6DFD<=6 AND PAC6DPAI<=9 AND SDC6GCB<=4 AND SDC6GRAC<=2 AND SMC6_3<=996 AND SMC6_4<=996 AND DGC6_1G<=6 AND CCC6_1H<=2 AND DGK6_1<=6 AND DHC6_OWN<=2 AND DHC6GBED<=5 AND DHC6GMAR<=3 AND DHC6GECF<=99 AND DHC6DLVG<=10; if PRC6 CUR=10 THEN NFLD=1; ELSE NFLD=0; IF PRC6_CUR=11 THEN PEI=1; ELSE PEI=0; /*IF PRC6_CUR=12 THEN NS=1; ELSE NS =0; */ IF PRC6_CUR=13 THEN NB=1; ELSE NB=0; IF PRC6_CUR=24 THEN QUE=1; ELSE QUE=0; IF PRC6_CUR=35 THEN ONT=1; ELSE ONT=0; IF PRC6 CUR=46 THEN MB=1; ELSE MB=0; IF PRC6_CUR=47 THEN SASK=1; ELSE SASK=0; IF PRC6_CUR=48 THEN AB=1; ELSE AB=0; IF PRC6_CUR=59 THEN BC=1; ELSE BC=0; =1; ELSE GE36GU1 =0;TF GE36GURB= 1 THEN GE36GU1 IFGE36GURB= 2 THEN GE36GU2 =1; ELSE GE36GU2 =0; IF GE36GCMA= 462 THEN GE36GC1 =1; ELSE GE36GC1 =0; IF GE36GCMA= 933 THEN GE36GC2 =1; ELSE GE36GC2 =0;

IF	GE36GHLR=	461	THEN	GE36GH1	=1;	ELSE	GE36GH1	=0;
IF	GE36GHLR=	462	THEN	GE36GH2	=1;	ELSE	GE36GH2	=0;
IF	GE36GHLR=	463	THEN	GE36GH3	=1;	ELSE	GE36GH3	=0;
IF	GE36GHLR=	464	THEN	GE36GH4	=1;	ELSE	GE36GH4	=0;
IF	GE36GHLR=	465	THEN	GE36GH5	=1;	ELSE	GE36GH5	=0;
IF	GE36GHLR=	481	THEN	GE36GH6	=1;	ELSE	GE36GH6	=0;
IF	GE36GHLR=	482	THEN	GE36GH7	=1;	ELSE	GE36GH7	=0;
IF	GE36GHLR=	483	THEN	GE36GH8	=1;	ELSE	GE36GH8	=0;
IF	GE36GHLR=	484	THEN	GE36GH9	=1;	ELSE	GE36GH9	=0;
IF	GE36GHLR=	485	THEN	GE36GH10	=1;	ELSE	GE36GH10	=0;
IF	GE36GHLR=	3511	THEN	GE36GH11	=1;	ELSE	GE36GH11	=0;
IF	GE36GHLR=	3512	THEN	GE36GH12	=1;	ELSE	GE36GH12	=0;
IF	GE36GHLR=	3521	THEN	GE36GH12 GE36GH13	=1;	ELSE	GE36GH12 GE36GH13	=0; =0;
IF	GE36GHLR=	3522	THEN	GE36GH14	=1;		GE36GH13 GE36GH14	
						ELSE		=0;
IF	GE36GHLR=	3523	THEN	GE36GH15	=1;	ELSE	GE36GH15	=0;
IF	GE36GHLR=	3524	THEN	GE36GH16	=1;	ELSE	GE36GH16	=0;
IF	GE36GHLR=	3531	THEN	GE36GH17	=1;	ELSE	GE36GH17	=0;
IF	GE36GHLR=	3532	THEN	GE36GH18	=1;	ELSE	GE36GH18	=0;
IF	GE36GHLR=	3533	THEN	GE36GH19	=1;	ELSE	GE36GH19	=0;
IF	GE36GHLR=	3534	THEN	GE36GH20	=1;	ELSE	GE36GH20	=0;
IF	GE36GHLR=	3541	THEN	GE36GH21	=1;	ELSE	GE36GH21	=0;
IF	GE36GHLR=	3542	THEN	GE36GH22	=1;	ELSE	GE36GH22	=0;
IF	GE36GHLR=	3543	THEN	GE36GH23	=1;	ELSE	GE36GH23	=0;
IF	GE36GHLR=	3551	THEN	GE36GH24	=1;	ELSE	GE36GH24	=0;
IF	GE36GHLR=	3552	THEN	GE36GH25	=1;	ELSE	GE36GH25	=0;
IF	GE36GHLR=	3561	THEN	GE36GH26	=1;	ELSE	GE36GH26	=0;
IF	GE36GHRO=	461	THEN	GE36G1	=1;	ELSE	GE36G1	=0;
IF	GE36GHRO=	462	THEN	GE36G2	=1;	ELSE	GE36G2	=0;
IF	GE36GHRO=	463	THEN	GE36G3	=1;	ELSE	GE36G3	=0;
IF	GE36GHRO=	464	THEN	GE36G4	=1;	ELSE	GE36G4	=0;
IF	GE36GHRO=	465	THEN	GE36G5	=1;	ELSE	GE36G5	=0;
IF	GE36GHRO=	481	THEN	GE36G6	=1;	ELSE	GE36G6	=0;
IF	GE36GHRO=	482	THEN	GE36G7	=1;	ELSE	GE36G7	=0;
IF	GE36GHRO=	483	THEN	GE36G8	=1;	ELSE	GE36G8	=0;
IF	GE36GHRO=	484	THEN	GE36G9	=1;	ELSE	GE36G9	=0;
IF	GE36GHRO=	485	THEN	GE36G10	=1;	ELSE	GE36G10	=0;
IF	GE36GHRO=	3511	THEN	GE36G11	=1;	ELSE	GE36G11	=0;
IF	GE36GHRO=	3512	THEN	GE36G12	=1; =1;	ELSE	GE36G12	=0; =0;
IF	GE36GHRO=	3512	THEN	GE36G12 GE36G13	=1;	ELSE	GE36G12 GE36G13	=0;
IF	GE36GHRO=	3521	THEN	GE36G14	=1; =1;		GE36G14	=0; =0;
IF	GE36GHRO=	3521				ELSE		
			THEN	GE36G15	=1;	ELSE	GE36G15	=0;
IF	GE36GHRO=	3523	THEN	GE36G16	=1;	ELSE	GE36G16	=0;
IF	GE36GHRO=	3524	THEN	GE36G17	=1;	ELSE	GE36G17	=0;
IF	GE36GHRO=	3525	THEN	GE36G18	=1;	ELSE	GE36G18	=0;
IF	GE36GHRO=	3526	THEN	GE36G19	=1;	ELSE	GE36G19	=0;
IF	GE36GHRO=	3527	THEN	GE36G20	=1;	ELSE	GE36G20	=0;
IF	GE36GHRO=	3531	THEN	GE36G21	=1;	ELSE	GE36G21	=0;
IF	GE36GHRO=	3532	THEN	GE36G22	=1;	ELSE	GE36G22	=0;
IF	GE36GHRO=	3533	THEN	GE36G23	=1;	ELSE	GE36G23	=0;
IF	GE36GHRO=	3534	THEN	GE36G24	=1;	ELSE	GE36G24	=0;
IF	GE36GHRO=	3536	THEN	GE36G25	=1;	ELSE	GE36G25	=0;
IF	GE36GHRO=	3541	THEN	GE36G26	=1;	ELSE	GE36G26	=0;
IF	GE36GHRO=	3542	THEN	GE36G27	=1;	ELSE	GE36G27	=0;
IF	GE36GHRO=	3543	THEN	GE36G28	=1;	ELSE	GE36G28	=0;
IF	GE36GHRO=	3544	THEN	GE36G29	=1;	ELSE	GE36G29	=0;

IF	GE36GHRO=	3551	THEN	GE36G30	=1;	ELSE	GE36G30	=0;
IF	GE36GHRO=	3552	THEN	GE36G31	=1;	ELSE	GE36G31	=0;
IF	GE36GHRO=	3553	THEN	GE36G32	=1;	ELSE	GE36G32	=0;
IF	GE36GHRO=	3561	THEN	GE36G33	=1;	ELSE	GE36G33	=0;
IF	0<=HWC6GBMI	<=57.9	THEN	HWC6GB1	=HWC6	GBMI;	ELSE HWC6G	В1
	=0;							
IF	HWC6GSW=	1	THEN	HWC6G1	=1;	ELSE	HWC6G1	=0;
IF	HWC6GSW=	2	THEN	HWC6G2	=1;	ELSE	HWC6G2	=0;
IF	HWC6GSW=	3	THEN	HWC6G3	=1;	ELSE	HWC6G3	=0;
IF	HWC6GSW=	4	THEN	HWC6G4	=1;	ELSE	HWC6G4	=0;
IF	HWS6_1=	1	THEN	HWS1 $=1;$	ELSE	HWS1	=0;	
IF	HWS6_1=	2	THEN	HWS2 =1;	ELSE	HWS2	=0;	
IF	$HWS6_1 =$	3	THEN	HWS3 =1;	ELSE	HWS3	=0;	
	—							
* IF	0<=HWS6_2KG	-185	THEN	HWS6KG1	=HWS6	280.	* ELSE	
τt.			TITT	HWSONGT	-11050	_210;		
	HWS6KG1	=0;						
	-							
IF	HWS6_5=	1	THEN	HWS61 =1;	ELSE	HWS61		
IF	HWS6_5=	2	THEN	HWS62 =1;	ELSE	HWS62	=0;	
IF	LFC6GO21=	1	THEN	LFC6G1	=1;	ELSE	LFC6G1	=0;
IF	LFC6GO21=	2	THEN	LFC6G2	=1;	ELSE	LFC6G2	=0;
IF	LFC6G021=	3	THEN	LFC6G3	=1;	ELSE	LFC6G3	=0;
					=1;			
IF	LFC6GO21=	4	THEN	LFC6G4		ELSE	LFC6G4	=0;
IF	LFC6GO21=	5	THEN	LFC6G5	=1;	ELSE	LFC6G5	=0;
IF	LFC6GO21=	6	THEN	LFC6G6	=1;	ELSE	LFC6G6	=0;
IF	LFC6GO21=	7	THEN	LFC6G7	=1;	ELSE	LFC6G7	=0;
IF	LFC6GO21=	8	THEN	LFC6G8	=1;	ELSE	LFC6G8	=0;
IF	LFC6GO21=	9	THEN	LFC6G9	=1;	ELSE	LFC6G9	=0;
IF	LFC6G021=	10	THEN	LFC6G10	=1;	ELSE	LFC6G10	=0;
IF	LFC6GO21=	11	THEN	LFC6G11	=1;	ELSE	LFC6G11	=0;
IF	LFC6GO21=	12	THEN	LFC6G12	=1;	ELSE	LFC6G12	=0;
IF	LFC6GO21=	13	THEN	LFC6G13	=1;	ELSE	LFC6G13	=0;
IF	LFC6GO21=	14	THEN	LFC6G14	=1;	ELSE	LFC6G14	=0;
IF	LFC6GO21=	15	THEN	LFC6G15	=1;	ELSE	LFC6G15	=0;
IF	LFC6GO21=	16	THEN	LFC6G16	=1;	ELSE	LFC6G16	=0;
IF	LFC6GO21=	17	THEN	LFC6G17	=1;	ELSE	LFC6G17	=0;
IF	LFC6G021=	18	THEN	LFC6G18	=1;	ELSE	LFC6G18	=0;
IF	LFC6G021=	19	THEN	LFC6G19	=1;	ELSE	LFC6G19	=0;
IF	LFC6G021=	20	THEN	LFC6G20	=1;	ELSE	LFC6G20	=0;
IF	LFC6GO21=	21	THEN	LFC6G21	=1;	ELSE	LFC6G21	=0;
IF	LFC6GI13=	96	THEN	LFC1 = 1;	ELSE	LFC1	=0;	
IF	LFC6GI13=	2	THEN	LFC2 $=1;$	ELSE	LFC2	=0;	
IF	LFC6GI13=	3	THEN	LFC3 =1;	ELSE	LFC3	=0;	
IF	LFC6GI13=	4	THEN	LFC4 $=1;$	ELSE	LFC4	=0;	
IF	LFC6GI13=	5	THEN	LFC5 = 1;	ELSE	LFC5	=0;	
IF	LFC6GI13=	6	THEN	LFC6 = 1;	ELSE	LFC6	=0;	
IF	LFC6GI13=	7	THEN	LFC7 =1;	ELSE	LFC7	=0;	
IF	LFC6GI13=	8	THEN	LFC8 =1;	ELSE	LFC8	=0;	
IF	LFC6GI13=	9	THEN	LFC9 =1;	ELSE	LFC9	=0;	
IF	LFC6GI13=	10	THEN	LFC10 =1;	ELSE	LFC10	=0;	
IF	LFC6GI13=	11	THEN	LFC11 =1;	ELSE	LFC11	=0;	
IF	LFC6GI13=	12	THEN	LFC12 =1;	ELSE	LFC12		
				,		010	-,	

IF	LFC6GI13=	13	THEN	LFC13	=1;	ELSE	LFC13	=0;		
IF	0<=PAC6DEE< =0;	=34.4	THEN	PAC6D	E1	=PAC6	DEE;	ELSE	PAC6D	E1
IF IF	PAC6DLEI=1 PAC6DLEI=2	THEN THEN	PAC6D PAC6D		=1; =1;	ELSE ELSE	PAC6DI PAC6DI		=0; =0;	
IF	0 <pac6dfm<= ELSE if PAC6 ELSE PAC</pac6dfm<= 	DFM=0		PAC6D AC6DFM		=PAC6 ;	DFM;			
IF	PAC6DFR=	1	THEN	PAC6D	FR1	=1;	ELSE	PAC6D	FR1	=0;
IF	PAC6DFR=	2	THEN	PAC6D		=1;	ELSE	PAC6D		=0;
IF	PAC6DFR=	3	THEN	PAC6D		=1;	ELSE	PAC6D		=0;
IF	PAC6DFD=	1	THEN	PAC6D	FD1	=1;	ELSE	PAC6D	FD1	=0;
IF	PAC6DFD=	2	THEN	PAC6D	FD2	=1;	ELSE	PAC6D	FD2	=0;
IF	PAC6DPAI=	1	THEN	PAC6D	PA1	=1;	ELSE	PAC6D	PA1	=0;
IF	PAC6DPAI=	2	THEN	PAC6D	PA2	=1;	ELSE	PAC6D	PA2	=0;
IF	PAC6DPAI=	3	THEN	PAC6D	PA3	=1;	ELSE	PAC6D	PA3	=0;
IF	SDC6GCB=	1	THEN	SDC6G	1	=1;	ELSE	SDC6G	1	=0;
IF	SDC6GCB=	2	THEN	SDC6G	2	=1;	ELSE	SDC6G	2	=0;
IF	SDC6GCB=	3	THEN	SDC6G	3	=1;	ELSE	SDC6G	3	=0;
/* IF */	SDC6GCB=	4	THEN	SDC6G	4	=1;	ELSE	SDC6G	4	=0;
IF	SDC6GRAC=	1	THEN	SDC6G	R1	=1;	ELSE	SDC6G	R1	=0;
/*IF	SDC6GRAC=	2	THEN	SDC6G		=1;	ELSE	SDC6G		=0;*/
IF	0<=SMC6_3<= SMC6_31	85 =100;		THEN	SMC6_	31	=SMC6	_3;	ELSE	
IF	SMC6_3=996	-100,	THEN	SMC6_	32	=0;	ELSE	SMC6_	32	=1;
IF	0<=SMC6_4<= SMC6_41	99 =0;		THEN	SMC6_	41	=SMC6	_4;	ELSE	
IF	DGC6_1G=	1	THEN	DGC6G	:1	=1;	ELSE	DGC6G	1	=0;
IF	DGC6_1G= DGC6_1G=	2	THEN	DGC 6G		=1;	ELSE	DGC6G		=0;
IF	ССС6_1н=	1	THEN	ССС6н	11	=1;	ELSE	ССС6н	1	=0;
/*IF */	CCC6_1H=	2	THEN	ССС6н	12	=1;	ELSE	ССС6Н	2	=0;
IF	DGK6_1=	1	THEN	DCK61	. =1;	ELSE	DGK61	=0.		
IF	DGK6_1=	2	THEN		=1;	ELSE	DGK61 DGK62			
/*IF	DHC6_OWN=	1	THEN	DHC6_	01	=1;	ELSE	DHC6_	01	=0;*/
IF	DHC6_OWN=	2	THEN	DHC6_		=1;	ELSE			=0;
*IF	DHC6GBED= =0;	0	THEN	DHC60	B1	=1;	*ELSE		DHC6G	B1
*IF	=0; DHC6GBED= =0;	1	THEN	DHC6G	B2	=1;	*ELSE		DHC6G	B2

*IF	DHC6GBED= =0;	2	THEN	DHC6GB3	=1;	*ELSE	DHC6G	B3
*IF	DHC6GBED= =0;	3	THEN	DHC6GB4	=1;	*ELSE	DHC6G	B4
*IF	DHC6GBED= =0;	4	THEN	DHC6GB5	=1;	*ELSE	DHC6G	в5
*IF	DHC6GBED= =0;	5	THEN	DHC6GB6	=1;	*ELSE	DHC6G	в6
IF	0<=DHC6GBEI =0;)<=5	THEN	DHC6GB1	=DHC6	GBED;	ELSE DHC6G	B1
/*IF	DHC6_SEX=	1	THEN	DHC6_S1	=1;	ELSE	DHC6_S1	=0;*/
IF	DHC6_SEX=	2	THEN	DHC6_S2	=1;	ELSE	DHC6_S2	=0;
/*IF	DHC6GMAR=	1	THEN	DHC6GM1	=1;	ELSE	DHC6GM1	=0;*/
/ II IF	DHC6GMAR=	2	THEN	DHC6GM2	=1;	ELSE	DHC6GM2	=0;
IF	DHC6GMAR=	3	THEN	DHC6GM3	=1;	ELSE	DHC6GM3	=0;
ΤĽ	DHCOGMAR-	5	TURN	DICOGINS	-1,	6036	DIICOGIAS	-0,
*IF	DHC6GHSZ= =0;	1	THEN	DHC6GH1	=1;	*ELSE	DHC6G	H1
*IF	DHC6GHSZ= =0;	2	THEN	DHC6GH2	=1;	*ELSE	DHC 6G	H2
*IF	DHC6GHSZ= =0;	3	THEN	DHC6GH3	=1;	*ELSE	DHC60	н3
*IF	DHC6GHSZ= =0;	4	THEN	DHC6GH4	=1;	*ELSE	DHC60	H4
*IF	DHC6GHSZ= =0;	5	THEN	DHC6GH5	=1;	*ELSE	DHC 6G	H5
IF	0<=DHC6GHS2	2<=5	THEN	DHC6GH	=1;	ELSE	DHC6GH=0;	
IF	DHC6GLE5≈	1	THEN	DHC6GLE1	=1;	ELSE	DHC6GLE1	=0;
/*IF	DHC6GLE5=	2	THEN	DHC6GLE2	=1;	ELSE	DHC6GLE2	=0;*/
IF	DHC6G611=	1	THEN	DHC6G61	=1;	ELSE	DHC6G61	=0;
/*IF	DHC6G611=	2	THEN	DHC6G62	=1;	ELSE	DHC6G62	=0;*/
IF	DHC6GECF=	1	THEN	DHC6GE1	=1;	ELSE	DHC6GE1	=0;
IF	DHC6GECF≈	2	THEN	DHC6GE2	=1;	ELSE	DHC6GE2	=0;
IF	DHC6GECF=	3	THEN	DHC6GE3	=1;	ELSE	DHC6GE3	=0;
IF	DHC6GECF=	4	THEN	DHC6GE4	=1;	ELSE	DHC6GE4	=0;
IF	DHC6GECF=	5	THEN	DHC6GE5	=1;	ELSE	DHC6GE5	=0;
IF	DHC6GECF=	6	THEN	DHC6GE6	=1;	ELSE	DHC6GE6	=0;
								=0;*/
/*IF	DHC6GECF≈	7	THEN	DHC6GE7	=1;	ELSE	DHC6GE7	
IF	DHC6GECF=	8	THEN	DHC6GE8	=1;	ELSE	DHC6GE8	=0;
IF	DHC6GECF=	9	THEN	DHC6GE9	=1;	ELSE	DHC6GE9	=0;
IF	DHC6DLVG=	1	THEN	DHC6DL1	=1;	ELSE	DHC6DL1	=0;
IF	DHC6DLVG=	2	THEN	DHC6DL2	=1;	ELSE	DHC6DL2	=0;
IF	DHC6DLVG=	3	THEN	DHC6DL3	=1;	ELSE	DHC6DL3	=0;
IF	DHC6DLVG=	4	THEN	DHC6DL4	=1;	ELSE	DHC6DL4	=0;
IF	DHC6DLVG=	4 5	THEN		=1; =1;	ELSE	DHC6DL5	=0; =0;
				DHC6DL5				
IF (*TD	DHC6DLVG=	6	THEN	DHC6DL6	=1;	ELSE	DHC6DL6	=0;
/*IF	DHC6DLVG=	7	THEN	DHC6DL7	=1;	ELSE	DHC6DL7	=0;*/
IF	DHC6DLVG=	8	THEN	DHC6DL8	=1;	ELSE	DHC6DL8	=0;
IF	DHC6DLVG=	9	THEN	DHC6DL9	=1;	ELSE	DHC6DL9	=0;
IF	DHC6DLVG=	10	THEN	DHC6DL10	=1;	ELSE	DHC6DL10	=0;
run;								

data lgtdata; set lgtdata1; IF 6<DHC6GAGE; run; data lgtdata; set lgtdata; mul=rantbl(0,0.333333,0.333333,0.333334); run; data a; set lgtdata; if mul=1; run; data b; set lgtdata; if mul=2; run; data c; set lgtdata; if mul=3; run; ***** proc logistic data =a DESCENDING; weight WT66; model CCC6 1C= NFLD PEI NB QUE ONT MB SASK AB BC GE36GU1-GE36GU2 HWC6GB1 HWC6G1-HWC6G4 HWS61-HWS62 LFC1-LFC13 PAC6DFM1 PAC6DPA1-PAC6DPA3 SDC6G1-SDC6G3 SMC6_31 SMC6_41 DHC6_02 DHC6GB1 DHC6_S2 DHC6GM2-DHC6GM3 DHC6G61 DHC6GE1-DHC6GE6 DHC6GE8-DHC6GE9 DHC6DL2-DHC6DL6 DHC6DL8-DHC6DL9 HWC6GHT DHC6GAGE /lackfit SELECTION=STEPWISE link=LOGIT; run; **** proc logistic data =b DESCENDING; weight WT66; model CCC6_1C= BC GE36GU2 HWC6G1 HWC6G2 LFC1LFC3 LFC7 LFC8 LFC13

PAC6DPA1 PAC6DPA2 SDC6G1 SDC6G2 DHC6_02 DHC6_S2 DHC6GM2 DHC6G61 DHC6GE4 DHC6GE5 HWC6GHT DHC6GAGE ; / outroc=roc1 roceps=1e-5; output out=outp p=p; run; ***** data out; set c; -2.5267 ee=EXP(-BC * 0.1365 * 0.2500 -GE36GU2 * 0.00062 -HWC6G1 * -HWC6G2 0.2273 * 0.3750 +LFC1 * -lfC3 0.2381 * 0.4545 -lfC7 * 0.2375 -lfC8 +LFC13 * 0.2692 * 0.0665 +PAC6DPA1 0.0357 * +PAC6DPA2 * +SDC6G1 0.9132 +SDC6G2 * 0.8882 * +DHC6_02 0.4687 0.2969 * +DHC6_S2 * -DHC6GM2 0.0262 * 0.00820 -DHC6G61 * +DHC6GE4 0.5963 * 0.0189 -DHC6GE5 -HWC6GHT * 0.00572 * -DHC6GAGE 0.0905); P=EE/(1+EE);if ccc6_lc=1 then asthma=1; else asthma=0; RUN; data out; SET OUT; keep asthma p; run; proc sort data=out; BY descending P; run; ******

```
#In Splus:
roc_function(out)
{out<-as.matrix(out)</pre>
 # se<-0
# sp1<-0
# n<-0
 # m1<-0
 # m2<-0
n<-dim(out)[1]</pre>
m1 < -sum(out[,2])
m2 < -n-m1
s3<-rep(0,n)
s4<-rep(0,n)
for (i in 1:n)
{
      s4[i]<-sum(out[1:i,2])
      s3[i]<-n-i-sum(out[i:n,2])
}
se<-s4/ml
sp1<-1-s3/m2
lines(spl,se)
}
#pp<-out[,1]</pre>
#p0<-out[1,1]</pre>
#for (i in 2:n)
#{
#
      if (p0-pp[i]>=0.001) p0<-pp[i]
#
             else {
              se[i]<-NA
#
#
              sp1[i]<-NA
#
             }
#}
#se<-se[!is.na(se)]</pre>
#sp1<-sp1[!is.na(sp1)]</pre>
plot(c(-0.0,1),c(-0.0,1),type='n', xlab='1-Specificity',
ylab='Sensitivity')
  lines(c(0,1),c(0,1))
roc(outic)
 lines(outib$.1MSPEC,outib$.SENSIT.,lty=3)
roc(outcc)
 lines(outcb$.1MSPEC,outcb$.SENSIT.,lty=3)
roc(outac)
 lines(outab$.1MSPEC,outab$.SENSIT.,lty=3)
```