TDLU Counts and PIH

Suzanne Dufault

April 9, 2019

Contents

1	Examining the Data	5
2	Zero-Inflated versus Hurdle Models2.1 Zero-Inflated2.2 Hurdle2.3 Follow-Up Questions	9
3	Testing for Zero-Inflation 3.1 Goodness-of-Fit Comparisons	10 10
4	Unadjusted Negative Binomial Model Results 4.1 Adjusted Negative Binomial Model Results	12
5	Tables and Figures for Paper	17
6	Supplemental	23

Abstract

Just as a brief background, the outcome, TDLU counts (terminal ductal lobular units), are counted by a specially trained pathologist, and represent the locations where breast cancer originates. The number of these units correlates with breast cancer risk, and our hypothesis is that women with PIH and the TT genotype will have lower TDLU counts, as this would support a possible mechanism for the association seen in prior studies. I am thinking you would do a similar type of genetic model as you did in the UK Biobank study to see if interaction (and possibly trend?) is present.

List of Tables

1 2	Summary statistics for whole cohort and broken down by PIH status [AMONG PIH POSITIVE WOMEN] Summary statistics by genotype	(
3	[AMONG PIH NEGATIVE WOMEN] Summary statistics by genotype	8
4	Number of non-zero TDLU counts	(
5	Testing for zero-inflation via model comparisons. The first three comparisons are nested models (with the latter nested in the former) and are compared via a likelihood ratio test. The last	
	comparison is between non-nested models and instead used the Vuong test for non-nested models.	10
6	Results of unadjusted negative binomial model. PIH compares non-PIH women (PIH = 0) to PIH women (PIH = 1). T alleles are treated as a factor with reference level of no t.alleles (t.alleles = 0). It appears that women with PIH are associated (not at 0.05 significance) with a lower incidence rate of TDLUs. Women with at least one T allele experience lower incidence rates of TDLUs (closer	
	to a 0.05 significance rate)	11
7	Results of unadjusted negative binomial model. PIH compares non-PIH women (PIH = 0) to PIH women (PIH = 1). T alleles are treated as a binary variable with the reference of no T alleles (T	
	alleles = 0). T alleles appear to have a protective effect (significant at 0.05 level) and PIH appears	1.1
8	protective (not significant at 0.05)	11
	history. Age at time of study, family history, and parity all seem to be significantly associated with	
	TDLU count (when adjusting for the rest of the model covariates). Increasing age has a protective	
	effect. Some family history has a harmful effect. Increasing parity has a harmful effect	11
9	Results of adjusted negative binomial model with interaction terms. PIH compares non-PIH women	
	(PIH = 0) to PIH women $(PIH = 1)$. T alleles are treated as a factor variable. The reference for	
	T alleles is no T alleles (T alleles = 0). The reference for family history is no family history. Age	
	at time of study and parity are both significantly associated with TDLU count (when adjusting for	
	the rest of the model covariates)	12
10	Results of adjusted negative binomial model without interaction term. PIH compares non-PIH	
	women (PIH = 0) to PIH women (PIH = 1). T alleles are treated a a binary variable. The	
	reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family	
	history. Age at time of study, family history, and parity all seem to be significantly associated with	
	TDLU count (when adjusting for the rest of the model covariates). Increasing age has a protective	
	effect. Some family history has a harmful effect. Increasing parity has a harmful effect	13
11	Results of adjusted negative binomial model with interaction terms. PIH compares non-PIH women (PIH = 0) to PIH women (PIH = 1). T alleles are treated as a binary variable. The reference	
	for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history.	
	Age at time of study and parity both seem to be significantly associated with TDLU count (when	16
10	adjusting for the rest of the model covariates)	13
12	[AMONG PIH POSITIVE WOMEN ONLY] Results of adjusted negative binomial model. T alleles are treated as a factor variable. The reference for T alleles is no T alleles (T alleles = 0). The	
	reference for family history is no family history. Among PIH positive women, having two T alleles	
	versus no T alleles appears protective (significant at 0.05 level). Age at first birth is approaching	
	significance (protective). Increasing age at study entry appears to be protective (significant at 0.05	
	level)	14
13	[AMONG PIH POSITIVE WOMEN ONLY] Results of adjusted negative binomial model. T alleles	1.
10	are treated as a binary variable. The reference for T alleles is no T alleles (T alleles = 0). The	
	reference for family history is no family history. Among PIH positive women, having two T alleles	
	versus no T alleles appears protective (approaching significance at 0.05 level). Age at first birth	
	is approaching significance (protective). Increasing age at study entry appears to be protective	
	(significant at 0.05 level)	14
14	[AMONG PIH POSITIVE WOMEN ONLY] Results of adjusted negative binomial model. T alleles	
	are treated linearly (e.g. as a linear trend). The reference for family history is no family history.	
	Among PIH positive women, increasing the number of T alleles appears to have a protective associa-	
	tion (significant at 0.05 level). Age at first birth is approaching significance (protective). Increasing	
	age at study entry appears to be protective (significant at 0.05 level)	14
		_

15	[AMONG PIH POSITIVE WOMEN ONLY] Goodness of fit. As seen, the most flexible model (Model 1) has no significant improvement in goodness of fit over the less flexible models (binary categorization and linear trend). Further, the non-nested Vuong test between the binary and trend	
	models does not reject the null hypothesis that the models are indistinguishable. This provides	
	evidence that either the binary or the linear model provide a sufficient fit to the data relative to	
16	the other models and both require the same degrees of freedom	15
	to have no assocation. Increasing age at study entry appears to be protective (significant at 0.05 level). Increasing parity appears to be harmful (significant at 0.05 level). Family history approaches	
17	significance (harmful)	15
	to have no assocation. Increasing age at study entry appears to be protective (significant at 0.05	
	level). Increasing parity appears to be harmful (significant at 0.05 level). Family history approaches	
	significance (harmful)	16
18	[AMONG PIH NEGATIVE WOMEN ONLY] Results of adjusted negative binomial model. T	
	alleles are treated as a linear variable. The reference for family history is no family history. Among	
	PIH negative women, T alleles appear to have no assocation. Increasing age at study entry appears	
	to be protective (significant at 0.05 level). Increasing parity appears to be harmful (significant at	1.0
19	0.05 level). Family history approaches significance (harmful)	16
LÐ	(Model 1) has no significant improvement in goodness of fit over the less flexible models (binary	
	categorization and linear trend). Further, the non-nested Vuong test between the binary and trend	
	models does not reject the null hypothesis that the models are indistinguishable. This provides	
	evidence that either the binary or the linear model provide a sufficient fit to the data relative to	
	the other models and both require the same degrees of freedom.	16
1	Summary statistics for whole cohort and broken down by PIH status	18
2	[AMONG PIH POSITIVE WOMEN] Summary statistics by genotype	19
3	[AMONG PIH NEGATIVE WOMEN] Summary statistics by genotype	20
2	Results of adjusted negative binomial model with interaction terms	21
3	Summary table of the adjusted T allele IRRs from the models stratified on HDP-status	22
S1	Goodness of fit test results	23
52	Complete adjusted negative binomial model with interaction terms	23
53	Complete adjusted negative binomial models stratified by HDP status	24
54	Complete adjusted negative binomial models stratified by HDP status	24
55	Complete adjusted negative binomial models stratified by HDP status with alleles treated linearly	
	for trend	24
56	Complete adjusted negative binomial models stratified by HDP status with alleles treated linearly	or
	ton tuon d	•) 🗆

List	αf	Figures
பக	$\mathbf{O}_{\mathbf{I}}$	riguics

1 Examining the Data

TABLE 1: Summary statistics for whole cohort and broken down by PIH status.

	Total (N = 191)	PIH: $0 \text{ (N} = 115)$	PIH: $1 \text{ (N} = 76)$
T Alleles			, ,
0 T Alleles	42 (21.99%)	24 (20.87%)	18 (23.68%)
1 T Alleles	98 (51.31%)	61 (53.04%)	37 (48.68%)
2 T Alleles	51 (26.70%)	30 (26.09%)	21 (27.63%)
Age at Entry to Study (years)		,	,
Mean	45.9	46.2	45.45
SD	10.61	10.45	10.89
Min	27	27	27
Max	66	66	66
BMI			
Mean	30.02	28.43	32.43
SD	8.01	7.38	8.36
Min	14.63	14.63	20.05
Max	70.6	58.8	70.6
Parity			
1 Child	43 (22.51%)	26 (22.61%)	17 (22.37%)
2 Children	100 (52.36%)	64 (55.65%)	36 (47.37%)
3+ Children	48 (25.13%)	25 (21.74%)	23 (30.26%)
Age at First Birth			,
Mean	27.02	27.32	26.57
SD	5.07	4.9	5.33
Min	15	19	15
Max	43	43	39
Age at Menarche (years)			
Mean	12.52	12.6	12.41
SD	1.51	1.47	1.58
Min	8	9	8
Max	17	17	16
Family History: 1st Degree Relative			
No	140 (73.30%)	85 (73.91%)	55 (72.37%)
Yes	51 (26.70%)	30 (26.09%)	21 (27.63%)
TDLUs			
Mean	11.01	11.75	9.88
SD	14.27	15.09	12.96
Min	0	0	0
Max	99	99	70

TABLE 2: [AMONG PIH POSITIVE WOMEN] Summary statistics by genotype.

	T alleles: $0 \text{ (N} = 18)$	T alleles: $1 \text{ (N} = 37)$	T alleles: $2 (N = 21)$
Age at Entry to Study (years)			
Mean	44	44.73	47.95
SD	10.81	10.45	11.8
Min	27	28	29
Max	61	63	66
BMI			
Mean	29.76	32.27	35.02
SD	6.36	9.16	7.94
Min	20.05	20.05	22.15
Max	47.81	70.6	53.17
Parity			
1 Child	5 (27.78%)	9 (24.32%)	3 (14.29%)
2 Children	9 (50.00%)	18 (48.65%)	9 (42.86%)
3+ Children	4 (22.22%)	10 (27.03%)	9 (42.86%)
Age at First Birth			
Mean	26.89	26.57	26.29
SD	5.89	5.18	5.35
Min	17	15	18
Max	38	39	36
Age at Menarche (years)			
Mean	12.67	12.14	12.67
SD	1.57	1.64	1.46
Min	9	8	11
Max	15	16	16
Family History: 1st Degree Relative			
No	11 (61.11%)	29 (78.38%)	15 (71.43%)
Yes	7 (38.89%)	8 (21.62%)	6 (28.57%)
TDLUs			
Mean	16.28	9.14	5.71
SD	17.85	12.22	5.81
Min	0	0	0
Max	70	47	22

TABLE 3: [AMONG PIH NEGATIVE WOMEN] Summary statistics by genotype.

	T alleles: $0 \text{ (N} = 24)$	T alleles: $1 \text{ (N} = 61)$	T alleles: $2 (N = 30)$
Age at Entry to Study (years)	,		
Mean	48.58	46.16	44.37
SD	10.45	10.04	11.25
Min	30	27	29
Max	63	65	66
BMI			
Mean	29.31	28.7	27.15
SD	9.03	6.62	7.48
Min	19.41	17.96	14.63
Max	51.98	47.53	58.8
Parity			
1 Child	1 (4.17%)	17 (27.87%)	8 (26.67%)
2 Children	18 (75.00%)	31 (50.82%)	15 (50.00%)
3+ Children	5 (20.83%)	13 (21.31%)	7 (23.33%)
Age at First Birth	,		
Mean	27.08	27.56	27.03
SD	5.06	5.14	4.37
Min	20	19	19
Max	38	43	40
Age at Menarche (years)			
Mean	12.5	12.25	13.4
SD	1.5	1.31	1.48
Min	9	9	11
Max	17	16	16
Family History: 1st Degree Relative			
No	18 (75.00%)	43 (70.49%)	24 (80.00%)
Yes	6 (25.00%)	18 (29.51%)	6 (20.00%)
TDLUs			
Mean	13.46	10.77	12.37
SD	22.71	12.83	11.85
Min	0	0	0
Max	99	59	45

TDLU_cat	n	percent
non-zero	169	88.48
zero	22	11.52

TABLE 4: Number of non-zero TDLU counts.

2 Zero-Inflated versus Hurdle Models

Given that 10% of our data is comprised of zeros, we want to consider a zero-inflated model alongside the standard Poisson and Negative Binomial regression fits.

2.1 Zero-Inflated

Zero-inflated models assume that the zeros arise from two sources: *structural* and *sampling*. The *structural* component is those not at risk (e.g. non-smokers smoke 0 cigarettes, non-fishers catch 0 fish, etc.). The *sampling* component is those who are at risk, but by the count process, sample 0 anyway (e.g. fishers can catch 0 fish, sexually active individual may have 0 risky encounters, etc.)

The zero-inflated Poisson distribution for participant i can be defined as: [Source: Rose et al.]

$$P(Y_i = y_i) = \begin{cases} p_i + (1 - p_i)e^{-\mu_i} & y_i = 0\\ (1 - p_i)\frac{e^{-\mu_i}\mu_i^{y_i}}{y_i!} & y_i > 0 \end{cases}$$
 (1)

where p_i is the probability of being an excess zero and is often modeled using logistic regression:

$$logit(p_i) = X\beta \tag{2}$$

"Interpretation of the ZIP model depends upon what is being modeled. For medical studies the zero-inflated portion can be thought of as the odds of moving from the non-risk to the at-risk group. Once in the at-risk group we can determine the expected number of events or the risk of an event for one group versus another group."

A zero-inflated negative binomial model can be parameterized similarly.

2.2 Hurdle

A hurdle model, on the other hand, assumes all excess zeros are structural (e.g. only non-smokers smoke 0 cigarettes.) As such, it has two parts: 1) a binary response model, 2) a truncated-at-zero count model.

A hurdle model can be expressed as:

$$P(Y=0) = f_1(0) = p (3)$$

$$P(Y=i) = (1-p)\frac{f_2(i)}{1-f_2(0)} = (1-p)f_2'(i) \qquad i > 0$$
(4)

for any probability density functions f_1 and f_2 corresponding to non-negative integers. f_1 is the hurdle and f_2 represents the count process.

"This allows us to interpret the positive outcomes (>0) that result from passing the zero hurdle (threshold). The hurdle portion of the two-part model estimates the probability that the threshold is crossed. Theoretically the threshold could be any value, but its usually taken as zero because this is most often meaningful in the context of the study objectives."

2.3 Follow-Up Questions

1. For an individual who is screened, are zero counts structural? In other words, at the time of screening do the zero counts arise strictly from those who do not have breast cancer? Or are we allowing the possibility that a woman with breast cancer could also have a TDLU count of 0?

3 Testing for Zero-Inflation

We'll do comparative model fitting of the following models:

- Poisson model
- Negative binomial model (NB)
- Zero-inflated Poisson (ZIP)
- Zero-inflated negative binomial (ZINB)
- Hurdle Model with Poisson counting process (HPois)
- Hurdle Model with negative binomial counting process (HNB)

Nested models directly compared via likelihood ratio test:

- Poisson is nested within the NB model.
- Poisson is nested within ZIP.
- NB is nested with ZINB.

The Vuong test for non-nested models will be used to distinguish between the previous set of models and the hurdle models.

3.1 Goodness-of-Fit Comparisons

Models: TDLUs \sim PIH + I(T Alleles = 1) + I(T Alleles = 2)

alternative.hypothesis	p.value
NB > Poisson	0.000
ZIP > Poisson	0.000
ZINB > NB	0.660
NB > HNB	0.046

TABLE 5: Testing for zero-inflation via model comparisons. The first three comparisons are nested models (with the latter nested in the former) and are compared via a likelihood ratio test. The last comparison is between non-nested models and instead used the Vuong test for non-nested models.

According to this test, the standard negative binomial model fits better or just as well as any other model. We'll proceed with the negative binomial model.

4 Unadjusted Negative Binomial Model Results

	Coefficient	Standard Error	Z Value	p-value	IRR	CI.lb	CI.ub
(Intercept)	2.803	0.200	14.014	0.000	16.487	11.141	24.400
PIH	-0.234	0.179	-1.306	0.191	0.791	0.557	1.124
t.alleles1	-0.399	0.222	-1.800	0.072	0.671	0.434	1.036
t.alleles2	-0.473	0.251	-1.884	0.060	0.623	0.381	1.019

TABLE 6: Results of unadjusted negative binomial model. PIH compares non-PIH women (PIH = 0) to PIH women (PIH = 1). T alleles are treated as a factor with reference level of no t.alleles (t.alleles = 0). It appears that women with PIH are associated (not at 0.05 significance) with a lower incidence rate of TDLUs. Women with at least one T allele experience lower incidence rates of TDLUs (closer to a 0.05 significance rate).

	Coefficient	Standard Error	Z Value	p-value	IRR	CI.lb	CI.ub
(Intercept)	2.80	0.20	14.00	0.00	16.44	11.11	24.34
PIH	-0.23	0.18	-1.28	0.20	0.80	0.56	1.13
'T alleles'one or more	-0.42	0.21	-2.01	0.04	0.65	0.43	0.99

TABLE 7: Results of unadjusted negative binomial model. PIH compares non-PIH women (PIH = 0) to PIH women (PIH = 1). T alleles are treated as a binary variable with the reference of no T alleles (T alleles = 0). T alleles appear to have a protective effect (significant at 0.05 level) and PIH appears protective (not significant at 0.05).

A likelihood ratio test between these two unadjusted models shows that the binary treatment of the T alleles variable is no worse of a fit than the model comparing 0, 1, and 2 T alleles (p-value: 0.725).

4.1 Adjusted Negative Binomial Model Results

For the covariates for the smaller dataset, I would leave BMI, age first birth, and parity as they are. Age needs to be added, and please remove menarche. We do need to keep family history, but do you think we can combine them into one variable, such that if they answered yes to any of the questions we consider them a yes, otherwise a no? That may simplify the model.

	Coefficient	Standard Error	Z Value	p-value	IRR	CI.lb	CI.ub
(Intercept)	5.351	1.121	4.775	0.000	210.924	23.448	1897.319
PIH	-0.134	0.171	-0.787	0.432	0.874	0.626	1.222
'T alleles'1	-0.178	0.205	-0.865	0.387	0.837	0.560	1.252
'T alleles'2	-0.200	0.234	-0.854	0.393	0.819	0.517	1.296
AFB	0.006	0.017	0.345	0.730	1.006	0.973	1.040
Age	-0.044	0.008	-5.602	0.000	0.957	0.943	0.972
'Family Hist'some family history	0.378	0.185	2.043	0.041	1.460	1.016	2.099
BMI	-0.022	0.011	-1.987	0.047	0.978	0.957	1.000
Parity	0.305	0.106	2.870	0.004	1.357	1.102	1.671
Menarche	-0.091	0.058	-1.570	0.116	0.913	0.815	1.023

TABLE 8: Results of adjusted negative binomial model without interaction term. PIH compares non-PIH women (PIH = 0) to PIH women (PIH = 1). T alleles are treated as a factor variable. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history. Age at time of study, family history, and parity all seem to be significantly associated with TDLU count (when adjusting for the rest of the model covariates). Increasing age has a protective effect. Some family history has a harmful effect. Increasing parity has a harmful effect.

	Coefficient	Standard Error	Z Value	p-value	IRR	CI.lb	CI.ub
(Intercept)	5.056	1.135	4.453	0.000	156.895	16.954	1451.892
PIH	0.208	0.345	0.604	0.546	1.231	0.627	2.419
'T alleles'1	-0.027	0.268	-0.102	0.918	0.973	0.575	1.645
'T alleles'2	0.099	0.307	0.322	0.747	1.104	0.605	2.016
AFB	0.008	0.017	0.498	0.619	1.008	0.975	1.043
Age	-0.041	0.008	-5.232	0.000	0.960	0.945	0.975
'Family Hist'some family history	0.372	0.185	2.008	0.045	1.451	1.009	2.086
BMI	-0.022	0.011	-1.918	0.055	0.979	0.957	1.000
Parity	0.334	0.107	3.132	0.002	1.396	1.133	1.720
Menarche	-0.102	0.058	-1.763	0.078	0.903	0.806	1.011
PIH:'T alleles'1	-0.310	0.417	-0.743	0.457	0.734	0.324	1.661
PIH:'T alleles'2	-0.740	0.482	-1.536	0.124	0.477	0.185	1.227

TABLE 9: Results of adjusted negative binomial model with interaction terms. PIH compares non-PIH women (PIH = 0) to PIH women (PIH = 1). T alleles are treated as a factor variable. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history. Age at time of study and parity are both significantly associated with TDLU count (when adjusting for the rest of the model covariates).

4.1.1 Using Delta Method for Covariate Combinations

```
broom::tidy(m_nb_adj_int_centered)
## # A tibble: 12 x 5
## term
                                   estimate std.error statistic p.value
##
   <chr>
                                    <dbl>
## 1 (Intercept)
                                   1.85
                                           0.494
                                                       3.74 1.83e-4
                                   0.208
## 2 PIH
                                            0.345
                                                       0.604 5.46e-1
## 3 `T alleles`1
                                   -0.0275 0.268 -0.102 9.18e-1
## 4 `T alleles`2
                                   0.0990 0.307
                                                      0.322 7.47e-1
                                   0.00845 0.0170
                                                       0.498 6.19e-1
## 5 AFB
                                   -0.0408 0.00779 -5.23
## 6 Age
                                                                1.68e-7
                                                       2.01
## 7 `Family Hist`some family history 0.372
                                             0.185
                                                                4.47e-2
                                   -0.0215 0.0112 -1.92
                                                              5.51e-2
## 8 BMI
                                          0.107
                                                       3.13
## 9 Parity
                                    0.334
                                                                1.73e-3
## 10 Menarche
                                   -0.102 0.0578
                                                      -1.76
                                                             7.79e-2
## 11 PIH: T alleles 1
                                   -0.310 0.417
                                                      -0.743 4.57e-1
## 12 PIH: T alleles 2
                                   -0.740
                                            0.482 -1.54 1.24e-1
new_names <- str_replace(names(m_nb_adj_int_centered$coef), ":", "_")</pre>
new_names <- str_replace_all(new_names, "'", "")</pre>
new_names <- str_replace_all(new_names, " ", "")</pre>
# Comparing TT to GG for HDP+
comp_TT_GG_HDPplus <- deltaMethod(m_nb_adj_int_centered,</pre>
           "Talleles2 + PIH_Talleles2",
           parameterNames = new_names)
2*pnorm(comp_TT_GG_HDPplus$Estimate/comp_TT_GG_HDPplus$SE, mean = 0, sd = 1)
## [1] 0.07988333
# Comparing GT to GG for HDP+
comp_GT_GG_HDPplus <- deltaMethod(m_nb_adj_int_centered,</pre>
                              "Talleles1 + PIH_Talleles1",
                              parameterNames = new_names)
2*pnorm(comp_GT_GG_HDPplus$Estimate/comp_GT_GG_HDPplus$SE, mean = 0, sd = 1)
## [1] 0.2884442
# Comparing TT to GG for HDP-
broom::tidy(m_nb_adj_int_centered) %>%
filter(term == "`T alleles`2")
```

```
## # A tibble: 1 x 5
##
   term
          estimate std.error statistic p.value
##
                  <dbl>
                            <dbl>
                                      <dbl>
    <chr>
                                              <dbl>
## 1 `T alleles`2 0.0990
                             0.307
                                      0.322
                                              0.747
# Comparing GT to GG for HDP-
broom::tidy(m_nb_adj_int_centered) %>%
 filter(term == "`T alleles`1")
## # A tibble: 1 x 5
    term
                estimate std.error statistic p.value
##
    <chr>
                 <dbl> <dbl>
                                      <dbl>
                                             0.918
## 1 `T alleles`1 -0.0275 0.268 -0.102
```

	Coefficient	Standard Error	Z Value	p-value	IRR	CI.lb	CI.ub
(Intercept)	5.381	1.091	4.933	0.000	217.278	25.615	1843.072
PIH	-0.134	0.171	-0.783	0.434	0.875	0.626	1.222
'T alleles'one or more	-0.185	0.194	-0.954	0.340	0.831	0.569	1.215
AFB	0.006	0.017	0.348	0.728	1.006	0.973	1.040
Age	-0.044	0.008	-5.606	0.000	0.957	0.943	0.972
'Family Hist'some family history	0.381	0.185	2.056	0.040	1.463	1.018	2.103
BMI	-0.022	0.011	-2.009	0.045	0.978	0.957	0.999
Parity	0.303	0.106	2.871	0.004	1.354	1.101	1.665
Menarche	-0.093	0.056	-1.662	0.097	0.911	0.817	1.017

TABLE 10: Results of adjusted negative binomial model without interaction term. PIH compares non-PIH women (PIH = 0) to PIH women (PIH = 1). T alleles are treated a a binary variable. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history. Age at time of study, family history, and parity all seem to be significantly associated with TDLU count (when adjusting for the rest of the model covariates). Increasing age has a protective effect. Some family history has a harmful effect. Increasing parity has a harmful effect.

	Coefficient	Standard Error	Z Value	p-value	IRR	CI.lb	CI.ub
(Intercept)	5.090	1.108	4.594	0.000	162.368	18.508	1424.405
PIH	0.197	0.346	0.570	0.569	1.218	0.619	2.397
'T alleles'one or more	0.005	0.256	0.021	0.983	1.005	0.609	1.660
AFB	0.009	0.017	0.523	0.601	1.009	0.976	1.043
Age	-0.043	0.008	-5.464	0.000	0.958	0.944	0.973
'Family Hist'some family history	0.349	0.185	1.883	0.060	1.418	0.986	2.040
BMI	-0.021	0.011	-1.863	0.062	0.979	0.958	1.001
Parity	0.322	0.106	3.040	0.002	1.379	1.121	1.697
Menarche	-0.098	0.056	-1.752	0.080	0.907	0.813	1.012
PIH:'T alleles'one or more	-0.440	0.399	-1.103	0.270	0.644	0.295	1.407

TABLE 11: Results of adjusted negative binomial model with interaction terms. PIH compares non-PIH women (PIH = 0) to PIH women (PIH = 1). T alleles are treated as a binary variable. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history. Age at time of study and parity both seem to be significantly associated with TDLU count (when adjusting for the rest of the model covariates).

Using a likelihood ratio test to compare the models (without interaction terms) with 0, 1, and 2 T alleles versus the model with binary classification of one or more T alleles, we obtain a p-value of 0.913, indicating that the model with more levels is not a better fit than the one with fewer. We can use the model with the binary treatment of T alleles.

4.2 Adjusted Stratified Models

	Coefficient	Standard Error	Z Value	p-value	IRR	CI.lb	CI.ub
(Intercept)	8.545	1.887	4.528	0.000	5142.603	127.282	207778.285
'T alleles'1	-0.500	0.326	-1.536	0.124	0.606	0.320	1.148
'T alleles'2	-0.760	0.386	-1.967	0.049	0.468	0.219	0.997
AFB	-0.045	0.028	-1.628	0.104	0.956	0.905	1.009
Age	-0.023	0.013	-1.793	0.073	0.978	0.954	1.002
'Family Hist'some family history	0.071	0.303	0.235	0.814	1.074	0.593	1.946
BMI	-0.027	0.018	-1.522	0.128	0.973	0.940	1.008
Parity	-0.048	0.179	-0.267	0.790	0.953	0.671	1.354
Menarche	-0.224	0.095	-2.360	0.018	0.799	0.663	0.963

TABLE 12: [AMONG PIH POSITIVE WOMEN ONLY] Results of adjusted negative binomial model. T alleles are treated as a factor variable. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history. Among PIH positive women, having two T alleles versus no T alleles appears protective (significant at 0.05 level). Age at first birth is approaching significance (protective). Increasing age at study entry appears to be protective (significant at 0.05 level).

	Coefficient	Standard Error	Z Value	p-value	IRR	CI.lb	CI.ub
(Intercept)	8.805	1.791	4.917	0.000	6665.206	199.293	222912.492
'T alleles'one or more	-0.576	0.310	-1.857	0.063	0.562	0.306	1.033
AFB	-0.044	0.028	-1.585	0.113	0.957	0.906	1.011
Age	-0.025	0.013	-1.970	0.049	0.975	0.952	1.000
'Family Hist'some family history	0.062	0.304	0.204	0.839	1.064	0.587	1.929
BMI	-0.028	0.017	-1.609	0.108	0.972	0.940	1.006
Parity	-0.078	0.175	-0.444	0.657	0.925	0.657	1.303
Menarche	-0.233	0.092	-2.526	0.012	0.792	0.661	0.949

TABLE 13: [AMONG PIH POSITIVE WOMEN ONLY] Results of adjusted negative binomial model. T alleles are treated as a binary variable. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history. Among PIH positive women, having two T alleles versus no T alleles appears protective (approaching significance at 0.05 level). Age at first birth is approaching significance (protective). Increasing age at study entry appears to be protective (significant at 0.05 level).

	Coefficient	Standard Error	Z Value	p-value	IRR	CI.lb	CI.ub
(Intercept)	8.415	1.812	4.645	0.000	4516.286	129.566	157424.573
'T alleles'	-0.380	0.194	-1.960	0.050	0.684	0.468	1.000
AFB	-0.046	0.028	-1.645	0.100	0.955	0.904	1.009
Age	-0.022	0.013	-1.767	0.077	0.978	0.954	1.002
'Family Hist'some family history	0.103	0.300	0.342	0.733	1.108	0.615	1.996
BMI	-0.028	0.018	-1.560	0.119	0.973	0.939	1.007
Parity	-0.032	0.178	-0.182	0.856	0.968	0.684	1.371
Menarche	-0.220	0.093	-2.375	0.018	0.802	0.669	0.962

TABLE 14: [AMONG PIH POSITIVE WOMEN ONLY] Results of adjusted negative binomial model. T alleles are treated linearly (e.g. as a linear trend). The reference for family history is no family history. Among PIH positive women, increasing the number of T alleles appears to have a protective association (significant at 0.05 level). Age at first birth is approaching significance (protective). Increasing age at study entry appears to be protective (significant at 0.05 level).

Tests can be used to compare these three models with respect to goodness of fit. Specifically, we have three models:

Model 1 Treating T alleles as a factor. This is the most flexible model and it allows the effect of one T allele to differ from the effect of two T alleles.

Model 2 Treating T alleles as a binary variable. This essentially assumes a plateau effect where having one T allele is the same as having two T alleles.

Model 3 Treating T alleles as a linear variable. This enforces a linear dose-response relationship between the number of T alleles and the log count of TDLUs.

Model Comparison	χ^2 p-value
Model 1 v Model 2 (nested, LRT)	0.451
Model 1 v Model 3 (nested, LRT)	0.664
Model 2 v Model 3 (non-nested Vuong Test)	0.337

TABLE 15: [AMONG PIH POSITIVE WOMEN ONLY] Goodness of fit. As seen, the most flexible model (Model 1) has no significant improvement in goodness of fit over the less flexible models (binary categorization and linear trend). Further, the non-nested Vuong test between the binary and trend models does not reject the null hypothesis that the models are indistinguishable. This provides evidence that either the binary or the linear model provide a sufficient fit to the data relative to the other models and both require the same degrees of freedom.

	Coefficient	Standard Error	Z Value	p-value	IRR	CI.lb	CI.ub
(Intercept)	3.789	1.393	2.721	0.007	44.219	2.885	677.757
'T alleles'1	0.013	0.260	0.052	0.959	1.014	0.608	1.689
'T alleles'2	0.080	0.302	0.264	0.792	1.083	0.600	1.955
AFB	0.022	0.021	1.022	0.307	1.022	0.980	1.065
Age	-0.043	0.010	-4.302	0.000	0.958	0.940	0.977
'Family Hist'some family history	0.441	0.229	1.924	0.054	1.554	0.992	2.435
BMI	-0.020	0.014	-1.369	0.171	0.980	0.953	1.009
Parity	0.398	0.131	3.028	0.002	1.489	1.151	1.927
Menarche	-0.041	0.073	-0.556	0.578	0.960	0.831	1.109

TABLE 16: [AMONG PIH NEGATIVE WOMEN ONLY] Results of adjusted negative binomial model. T alleles are treated as a factor. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history. Among PIH negative women, T alleles appear to have no association. Increasing age at study entry appears to be protective (significant at 0.05 level). Increasing parity appears to be harmful (significant at 0.05 level). Family history approaches significance (harmful).

	Coefficient	Standard Error	Z Value	p-value	IRR	CI.lb	CI.ub
(Intercept)	3.699	1.371	2.699	0.007	40.419	2.752	593.541
'T alleles'one or more	0.035	0.248	0.142	0.887	1.036	0.637	1.684
AFB	0.022	0.021	1.031	0.303	1.022	0.981	1.065
Age	-0.043	0.010	-4.322	0.000	0.958	0.940	0.977
'Family Hist'some family history	0.427	0.228	1.872	0.061	1.533	0.980	2.397
BMI	-0.019	0.014	-1.332	0.183	0.981	0.953	1.009
Parity	0.401	0.132	3.052	0.002	1.494	1.154	1.933
Menarche	-0.035	0.070	-0.500	0.617	0.966	0.842	1.107

TABLE 17: [AMONG PIH NEGATIVE WOMEN ONLY] Results of adjusted negative binomial model. T alleles are treated as binary. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history. Among PIH negative women, T alleles appear to have no association. Increasing age at study entry appears to be protective (significant at 0.05 level). Increasing parity appears to be harmful (significant at 0.05 level). Family history approaches significance (harmful).

	Coefficient	Standard Error	Z Value	p-value	IRR	CI.lb	CI.ub
(Intercept)	3.737	1.346	2.776	0.006	41.951	2.999	586.751
'T alleles'	0.041	0.151	0.270	0.787	1.042	0.775	1.400
AFB	0.022	0.021	1.031	0.303	1.022	0.981	1.065
Age	-0.043	0.010	-4.308	0.000	0.958	0.940	0.977
'Family Hist'some family history	0.436	0.228	1.911	0.056	1.547	0.989	2.419
BMI	-0.020	0.014	-1.348	0.178	0.981	0.953	1.009
Parity	0.402	0.131	3.071	0.002	1.495	1.157	1.933
Menarche	-0.039	0.072	-0.550	0.582	0.961	0.836	1.106

TABLE 18: [AMONG PIH NEGATIVE WOMEN ONLY] Results of adjusted negative binomial model. T alleles are treated as a linear variable. The reference for family history is no family history. Among PIH negative women, T alleles appear to have no assocation. Increasing age at study entry appears to be protective (significant at 0.05 level). Increasing parity appears to be harmful (significant at 0.05 level). Family history approaches significance (harmful).

Again, tests can be used to compare these three models with respect to goodness of fit. Specifically, we have three models:

Model 1 Treating T alleles as a factor. This is the most flexible model and it allows the effect of one T allele to differ from the effect of two T alleles.

Model 2 Treating T alleles as a binary variable. This essentially assumes a plateau effect where having one T allele is the same as having two T alleles.

Model 3 Treating T alleles as a linear variable. This enforces a linear dose-response relationship between the number of T alleles and the log count of TDLUs.

Model Comparison	χ^2 p-value
Model 1 v Model 2 (nested, LRT)	0.796
Model 1 v Model 3 (nested, LRT)	0.902
Model 2 v Model 3 (non-nested Vuong Test)	0.466

TABLE 19: [AMONG PIH NEGATIVE WOMEN ONLY] Goodness of fit. As seen, the most flexible model (Model 1) has no significant improvement in goodness of fit over the less flexible models (binary categorization and linear trend). Further, the non-nested Vuong test between the binary and trend models does not reject the null hypothesis that the models are indistinguishable. This provides evidence that either the binary or the linear model provide a sufficient fit to the data relative to the other models and both require the same degrees of freedom.

5 Tables and Figures for Paper

TABLE 1: Summary statistics for whole cohort and broken down by PIH status.

		PIH: 0 (N = 115)	PIH: $1 (N = 76)$
T Alleles			
0 T Alleles	42 (21.99%)	24 (20.87%)	18 (23.68%)
1 T Alleles	98 (51.31%)	61 (53.04%)	37 (48.68%)
2 T Alleles	51 (26.70%)	30 (26.09%)	21 (27.63%)
Age at Entry to Study (years)		,	, ,
Mean	45.9	46.2	45.45
SD	10.61	10.45	10.89
Min	27	27	27
Max	66	66	66
BMI			
Mean	30.02	28.43	32.43
SD	8.01	7.38	8.36
Min	14.63	14.63	20.05
Max	70.6	58.8	70.6
Parity			
1 Child	43 (22.51%)	26 (22.61%)	17 (22.37%)
2 Children	100 (52.36%)	64 (55.65%)	36 (47.37%)
3+ Children	48 (25.13%)	25 (21.74%)	23 (30.26%)
Age at First Birth			
Mean	27.02	27.32	26.57
SD	5.07	4.9	5.33
Min	15	19	15
Max	43	43	39
Age at Menarche (years)			
Mean	12.52	12.6	12.41
SD	1.51	1.47	1.58
Min	8	9	8
Max	17	17	16
Family History: 1st Degree Relative			
No	140 (73.30%)	85 (73.91%)	55 (72.37%)
Yes	51 (26.70%)	30 (26.09%)	21 (27.63%)
TDLUs			
Mean	11.01	11.75	9.88
SD	14.27	15.09	12.96
Min	0	0	0
Max	99	99	70

TABLE 2: [AMONG PIH POSITIVE WOMEN] Summary statistics by genotype.

	T alleles: $0 \text{ (N} = 18)$	T alleles: $1 \text{ (N} = 37)$	T alleles: $2 (N = 21)$
Age at Entry to Study (years)			
Mean	44	44.73	47.95
SD	10.81	10.45	11.8
Min	27	28	29
Max	61	63	66
BMI			
Mean	29.76	32.27	35.02
SD	6.36	9.16	7.94
Min	20.05	20.05	22.15
Max	47.81	70.6	53.17
Parity			
1 Child	5 (27.78%)	9 (24.32%)	3 (14.29%)
2 Children	9 (50.00%)	18 (48.65%)	9 (42.86%)
3+ Children	4 (22.22%)	10 (27.03%)	9 (42.86%)
Age at First Birth			
Mean	26.89	26.57	26.29
SD	5.89	5.18	5.35
Min	17	15	18
Max	38	39	36
Age at Menarche (years)			
Mean	12.67	12.14	12.67
SD	1.57	1.64	1.46
Min	9	8	11
Max	15	16	16
Family History: 1st Degree Relative			
No	11 (61.11%)	29 (78.38%)	15 (71.43%)
Yes	7 (38.89%)	8 (21.62%)	6 (28.57%)
TDLUs			
Mean	16.28	9.14	5.71
SD	17.85	12.22	5.81
Min	0	0	0
Max	70	47	22

TABLE 3: [AMONG PIH NEGATIVE WOMEN] Summary statistics by genotype.

	T alleles: $0 \text{ (N} = 24)$	T alleles: $1 \text{ (N} = 61)$	T alleles: $2 \text{ (N} = 30)$
Age at Entry to Study (years)			
Mean	48.58	46.16	44.37
SD	10.45	10.04	11.25
Min	30	27	29
Max	63	65	66
BMI			
Mean	29.31	28.7	27.15
SD	9.03	6.62	7.48
Min	19.41	17.96	14.63
Max	51.98	47.53	58.8
Parity			
1 Child	1 (4.17%)	17 (27.87%)	8 (26.67%)
2 Children	18 (75.00%)	31 (50.82%)	15 (50.00%)
3+ Children	5 (20.83%)	13 (21.31%)	7 (23.33%)
Age at First Birth			
Mean	27.08	27.56	27.03
SD	5.06	5.14	4.37
Min	20	19	19
Max	38	43	40
Age at Menarche (years)			
Mean	12.5	12.25	13.4
SD	1.5	1.31	1.48
Min	9	9	11
Max	17	16	16
Family History: 1st Degree Relative			
No	18 (75.00%)	43 (70.49%)	24 (80.00%)
Yes	6 (25.00%)	18 (29.51%)	6 (20.00%)
TDLUs			
Mean	13.46	10.77	12.37
SD	22.71	12.83	11.85
Min	0	0	0
Max	99	59	45

TABLE 2: Results of adjusted negative binomial model with interaction terms.

	Coefficient	Standard Error	Z Value	p-value	IRR (95% CI)
HDP	0.208	0.345	0.604	0.546	1.231 (0.627, 2.419)
T alleles = 1	-0.027	0.268	-0.102	0.918	$0.973 \ (0.575, 1.645)$
T alleles $= 2$	0.099	0.307	0.322	0.747	$1.104\ (0.605,\ 2.016)$
$HDP \times T$ alleles = 1	-0.310	0.417	-0.743	0.457	$0.734\ (0.324,\ 1.661)$
$HDP \times T$ alleles = 2	-0.740	0.482	-1.536	0.124	$0.477 \ (0.185, \ 1.227)$

HDP compares HDP-positive women to HDP-negative women. T alleles are treated as a factor variable. The reference for T alleles is no T alleles (T alleles = 0). These results are adjusted for family history, age at biopsy, parity, age at menarche, age at first birth, and BMI. Full model covariates can be found in the supplemental material, Table S2.

TABLE 3: Summary table of the adjusted T allele IRRs from the models stratified on HDP-status.

	HDP-Negative		HDP-Positiv	e
	IRR (95% CI)	p-value	IRR (95% CI)	p-value
Factor Model				
T Alleles = 0	1 (ref)		1 (ref)	
T Alleles = 1	1.014 (0.608, 1.689)	0.959	0.606 (0.320, 1.148)	0.124
T Alleles = 2	1.083 (0.600, 1.955)	0.792	$0.468\ (0.219,\ 0.997)$	0.049
Linear Trend Model	,		, , ,	
Δ T Alleles	$1.042\ (0.775,\ 1.400)$	0.787	$0.684\ (0.468,\ 1.000)$	0.050

These results are adjusted for family history, age at biopsy, parity, age at menarche, age at first birth, and BMI. Full model covariates can be found in the supplemental material, Table S3 and Table S4.

```
## PIH
## 0
       1
## 115 76
## T alleles
## 0 1 2
## 42 98 51
## Parity
## 1 2
         3
## 43 100 38
## Family Hist
## 0 1 2
## 140 40 10
             1
## # A tibble: 1 x 8
   afb.mean age.mean men.mean bmi.mean t.alleles.mode pih.mode parity.mode
##
      <dbl> <dbl> <dbl> <chr>
                                               <chr>
                                                       <chr>
## 1
       27.0 45.9
                      12.5
                             30.0 1
                                                0
## # ... with 1 more variable: family.mode <chr>
```

6 Supplemental

TABLE S1: Goodness of fit test results

Alternative Hypothesis	p-value
GLM-NB > Poisson	0.000
ZIP > Poisson	0.000
ZINB > GLM-NB	0.660
GLM-NB > HNB	0.046

 $\operatorname{GLM-NB}=\operatorname{generalized}$ linear model with negative binomial distribution

Poisson = generalized linear model with Poisson distribution

 ${\it ZIP}={\it zero-inflated}$ poisson model

ZINB = zero-inflated negative binomial model

HNB = negative binomial hurdle model.

TABLE S2: Complete adjusted negative binomial model with interaction terms

	Coefficient	Standard Error	Z Value	p-value	IRR (95% CI)
(Intercept)	1.849	0.494	3.741	0.000	6.352 (2.411, 16.735)
HDP	0.208	0.345	0.604	0.546	$1.231\ (0.627,\ 2.419)$
T Alleles = 1	-0.027	0.268	-0.102	0.918	$0.973\ (0.575,\ 1.645)$
T Alleles = 2	0.099	0.307	0.322	0.747	$1.104 \ (0.605, \ 2.016)$
AFB	0.008	0.017	0.498	0.619	$1.008 \ (0.975, \ 1.043)$
Age	-0.041	0.008	-5.232	0.000	$0.96 \ (0.945, \ 0.975)$
Family History	0.372	0.185	2.008	0.045	$1.451\ (1.009,\ 2.086)$
BMI	-0.022	0.011	-1.918	0.055	$0.979\ (0.957,\ 1.000)$
Parity	0.334	0.107	3.132	0.002	1.396 (1.133, 1.720)
Menarche	-0.102	0.058	-1.763	0.078	0.903 (0.806, 1.011)
$HDP \times T$ alleles = 1	-0.310	0.417	-0.743	0.457	$0.734\ (0.324,\ 1.661)$
$HDP \times T$ alleles = 2	-0.740	0.482	-1.536	0.124	$0.477 \ (0.185, \ 1.227)$

T alleles are treated as factor variables. HDP compares HDP+ to HDP- women. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history.

TABLE S3: Complete adjusted negative binomial models stratified by HDP status

	Coefficient	Standard Error	Z Value	p-value	IRR (95% CI)
(Intercept)	4.157	0.803	5.177	0.000	63.899 (13.242, 308.331)
T Alleles = 1	-0.500	0.326	-1.536	0.124	0.606 (0.320, 1.148)
T Alleles = 2	-0.760	0.386	-1.967	0.049	$0.468 \ (0.219, \ 0.997)$
AFB	-0.045	0.028	-1.628	0.104	$0.956 \ (0.905, \ 1.009)$
Age	-0.023	0.013	-1.793	0.073	$0.978 \; (0.954, 1.002)$
Family History	0.071	0.303	0.235	0.814	$1.074\ (0.593,\ 1.946)$
BMI	-0.027	0.018	-1.522	0.128	$0.973\ (0.940,\ 1.008)$
Parity	-0.048	0.179	-0.267	0.790	$0.953 \ (0.671, \ 1.354)$
Menarche	-0.224	0.095	-2.360	0.018	$0.799\ (0.663,\ 0.963)$

For HDP+ women only. T alleles are treated as factor variables. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history.

TABLE S4: Complete adjusted negative binomial models stratified by HDP status

	Coefficient	Standard Error	Z Value	p-value	IRR (95% CI)
(Intercept)	1.296	0.493	2.629	0.009	3.653 (1.390, 9.598)
T Alleles = 1	0.013	0.260	0.052	0.959	1.014 (0.608, 1.689)
T Alleles = 2	0.080	0.302	0.264	0.792	$1.083\ (0.600,\ 1.955)$
AFB	0.022	0.021	1.022	0.307	$1.022\ (0.980,\ 1.065)$
Age	-0.043	0.010	-4.302	0.000	$0.958 \ (0.940, \ 0.977)$
Family History	0.441	0.229	1.924	0.054	$1.554 \ (0.992, \ 2.435)$
BMI	-0.020	0.014	-1.369	0.171	$0.98 \ (0.953, \ 1.009)$
Parity	0.398	0.131	3.028	0.002	1.489 (1.151, 1.927)
Menarche	-0.041	0.073	-0.556	0.578	$0.96 \ (0.831, \ 1.109)$

For HDP- women only. T alleles are treated as factor variables. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history.

TABLE S5: Complete adjusted negative binomial models stratified by HDP status with alleles treated linearly for trend

	Coefficient	Standard Error	Z Value	p-value	IRR (95% CI)
(Intercept)	4.047	0.749	5.401	0.000	57.204 (13.174, 248.392)
T Alleles	-0.380	0.194	-1.960	0.050	0.684 (0.468, 1.000)
AFB	-0.046	0.028	-1.645	0.100	0.955 (0.904, 1.009)
Age	-0.022	0.013	-1.767	0.077	$0.978 \ (0.954, \ 1.002)$
Family History	0.103	0.300	0.342	0.733	$1.108 \ (0.615, \ 1.996)$
BMI	-0.028	0.018	-1.560	0.119	$0.973 \ (0.939, \ 1.007)$
Parity	-0.032	0.178	-0.182	0.856	$0.968 \ (0.684, \ 1.371)$
Menarche	-0.220	0.093	-2.375	0.018	$0.802\ (0.669,\ 0.962)$

For HDP+ women only. T alleles are treated linearly. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history.

TABLE S6: Complete adjusted negative binomial models stratified by HDP status with alleles treated linearly for trend

	Coefficient	Standard Error	Z Value	p-value	IRR (95% CI)
(Intercept)	1.266	0.455	2.786	0.005	3.548 (1.455, 8.648)
T Alleles	0.041	0.151	0.270	0.787	1.042 (0.775, 1.400)
AFB	0.022	0.021	1.031	0.303	1.022 (0.981, 1.065)
Age	-0.043	0.010	-4.308	0.000	$0.958 \ (0.940, \ 0.977)$
Family History	0.436	0.228	1.911	0.056	1.547 (0.989, 2.419)
BMI	-0.020	0.014	-1.348	0.178	$0.981\ (0.953,\ 1.009)$
Parity	0.402	0.131	3.071	0.002	1.495 (1.157, 1.933)
Menarche	-0.039	0.072	-0.550	0.582	0.961 (0.836, 1.106)

For HDP- women only. T alleles are treated linearly. The reference for T alleles is no T alleles (T alleles = 0). The reference for family history is no family history.

Negative Binomial Model Expected versus Observed Counts

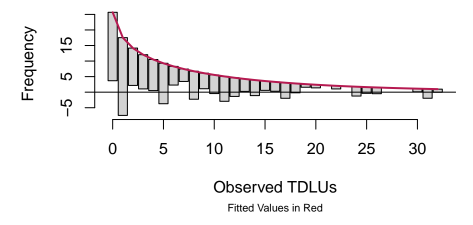


Figure 1: Rootogram. The bars represent the observed distribution of TDLU counts while the red line represents the model-fitted values. We see that we are overpredicting 0s, underpredicting 1s, overpredicting 2-4s, overpredicting 5s, etc.