Logistic Regression Models with GEE for the Secondary Outcomes in the VQI FBVAR Dataset

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Variables to adjust for

We used a correlation matrix to report the correlations between variables statistically. We also discussed with our collaborator on the choice of variables to adjust for. Here we did not report the correlation matrix.

- cluster on CENTERID
- adjust for AGECAT, GENDER, PREOP_SMOKING, PRIOR_AORSURG, PRIOR_CHF, PREOP_DIALYSIS
- adjust for PATHOLOGY, extent

In the model, we merge groups for the extent: merge "Juxtarenal AAA" with "Type 4 TAAA"; "Type 1 TAAA", "Type 2 TAAA", "Type 3 TAAA", with "Type 5 TAAA". Now extent is a binary variable, Juxtarenal or not.

• categorical secondary outcomes: POSTOP_CEREBROSX, POSTOP_SPINAL_ISCHEMIA, POSTOP_DIALYSIS, POSTOP_LOS, POSTOP_COMPLICATIONS, POSTOP_LEGEMBO, POSTOP_RESPIRATORY, RETX_R_RTOR, BRANCH_POST, TOTAL_LOS,ICUSTAY, POSTOP_INTISCH, POSTOP_GFR, POSTOP_HIGHCREAT

We use logistic regression with Generalized Estimating Equations to study categorical secondary outcomes

TOTAL_LOS: Length of stay in days between admission date and discharge date

	TOTAL_LOS Length of Stay is days				
Characteristic	OR^1	95% CI 1	p-value		
PRESENTATION					
Asymptomatic		_			
Symptomatic	5.14	4.11, 6.43	< 0.001		

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

	TOTAL	TOTAL_LOS Length of Stay in days			
Characteristic	OR^1	95% CI 1	p-value		
PRESENTATION					
Asymptomatic	_	_			
Symptomatic	3.79	2.99, 4.81	< 0.001		
AGECAT					
< 50	_	_			
>79	1.91	0.88, 4.15	0.10		
50-59	1.89	0.82, 4.36	0.13		
60-69	1.32	0.63, 2.77	0.47		
70-79	2.16	1.02, 4.55	0.043		
GENDER					
female		_			
male	0.62	0.51, 0.75	< 0.001		
PREOP_SMOKING					
No	_	_			
Yes	0.99	0.76, 1.29	0.94		
PRIOR_AORSURG					
Both		_			
Endo	1.59	0.93, 2.71	0.091		
None	1.08	0.65, 1.80	0.76		
Open	1.43	0.83, 2.47	0.20		
PRIOR_CHF					
No	_	_			
Yes	1.51	1.21, 1.89	< 0.001		
1 OR = Odds Ratio, CI = O	Confidence Inter	rval			

 $^{^{1}}OR = Odds Ratio, CI = Confidence Interval$

	TOTAL	_LOS Lengt days	h of Stay in
Characteristic	OR^1	95% CI ¹	p-value
PREOP_DIALYSIS			
No	_		
Yes	1.87	1.10, 3.16	0.020
PATHOLOGY			
Aneurysm	_		
Aneurysm from dissection	1.83	1.16, 2.87	0.009
Dissection	1.77	1.02, 3.06	0.041
PAU/IMH	1.94	0.90, 4.20	0.093
extent			
Juxtarenal	_	_	
No	2.10	1.72, 2.57	< 0.001

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

ICUSTAY: ICU Stay

		ICU Stay	7
Characteristic	OR^1	$95\%~\mathrm{CI}^1$	p-value
PRESENTATION			
Asymptomatic			
Symptomatic	3.11	2.43, 3.97	< 0.001

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

_	,	ICU Stay	Ţ
Characteristic	OR^1	95% CI ¹	p-value
PRESENTATION			
Asymptomatic	_	_	
Symptomatic	2.08	1.59, 2.71	< 0.001
AGECAT			
< 50	_	_	
>79	1.31	$0.51,\ 3.37$	0.58
50-59	1.30	0.47, 3.58	0.62
60-69	0.96	0.38, 2.42	0.93
70-79	1.28	0.50,3.25	0.61
GENDER			
female	_	_	
male	0.60	0.49, 0.74	< 0.001
PREOP_SMOKING			
No	_	_	
Yes	1.02	0.75, 1.38	0.92
PRIOR_AORSURG			
Both	_	_	
Endo	1.36	0.77, 2.41	0.29
None	0.75	0.43, 1.32	0.32
Open	1.31	0.72, 2.38	0.38
PRIOR_CHF			
No	_	_	
Yes	1.21	0.93, 1.57	0.17
PREOP_DIALYSIS			

 $^{^{1}\}mathrm{OR}=\mathrm{Odds}$ Ratio, CI = Confidence Interval

		ICU Stay	7
Characteristic	OR^1	95% CI ¹	p-value
No		_	
Yes	1.84	1.07, 3.19	0.028
PATHOLOGY			
Aneurysm	_	_	
Aneurysm from dissection	1.35	0.81,2.25	0.25
Dissection	1.72	0.93, 3.19	0.084
PAU/IMH	1.25	0.50, 3.12	0.63
extent			
Juxtarenal	_	_	
No	2.23	1.78, 2.80	< 0.001

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

Creatinine and GFR

 ${\tt POSTOP_HIGHCREAT: Highest\ Creatinine}$

	Highest Creatinine				
Characteristic	Beta	$95\%~\mathrm{CI}^1$	p-value		
PRESENTATION					
Asymptomatic		_			
Symptomatic	0.35	0.17, 0.52	< 0.001		

 $[\]overline{^{1}\text{CI} = \text{Confidence Interval}}$

	Highest Creatinine			
Characteristic	Beta	$95\%~\mathrm{CI}^1$	p-value	
PRESENTATION				
Asymptomatic	_			
Symptomatic	0.21	0.06, 0.37	0.006	
AGECAT				
< 50				
>79	-0.34	-0.89, 0.21	0.22	
50-59	-0.12	-0.72, 0.49	0.71	
60-69	-0.29	-0.85, 0.26	0.29	
70-79	-0.31	-0.86, 0.24	0.27	
GENDER				
female	_			
male	0.18	0.10,0.26	< 0.001	
PREOP_SMOKING				
No	_			
Yes	0.02	-0.08, 0.13	0.64	
PRIOR_AORSURG				
Both	_			
Endo	0.21	-0.06, 0.48	0.13	
None	0.11	-0.16, 0.39	0.41	
Open	0.05	-0.23, 0.33	0.72	
PRIOR_CHF				
No	_	_		
Yes	0.17	0.08,0.27	< 0.001	
1 CI = Confidence Interval				

	Highest Creatinine			
Characteristic	Beta	$95\%~\mathrm{CI}^1$	p-value	
PREOP_DIALYSIS				
No	_	_		
Yes	4.2	3.5, 5.0	< 0.001	
PATHOLOGY				
Aneurysm	_	_		
Aneurysm from dissection	-0.18	-0.43, 0.08	0.18	
Dissection	0.27	-0.13, 0.66	0.18	
PAU/IMH	0.19	-0.19, 0.56	0.33	
extent				
Juxtarenal	_	_		
No	0.07	-0.02, 0.16	0.15	

 $[\]overline{^{1}}$ CI = Confidence Interval

<code>GFRFIFTY: 1 - GFR</code> reduction is >=50%; 0 - GFR reduction is <50%

GFR reduction				
OR^1	95% CI 1	p-value		
	_			
1.57	1.08, 2.29	0.018		
		OR ¹ 95% CI ¹		

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

		GFR reduct	tion
Characteristic	OR^1	95% CI ¹	p-value
PRESENTATION			
Asymptomatic	_	_	
Symptomatic	1.35	0.90, 2.02	0.15
AGECAT			
< 50	_	_	
>79	0.72	0.15, 3.43	0.68
50-59	1.33	0.27,6.48	0.73
60-69	0.91	0.20,4.24	0.91
70-79	0.82	0.17, 3.87	0.81
1			

 $^{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

	GFR reduction			
Characteristic	OR^1	$95\%~\mathrm{CI}^1$	p-value	
GENDER				
female				
male	0.67	0.50, 0.89	0.006	
PREOP_SMOKING				
No	_	_		
Yes	1.27	0.81, 2.00	0.29	
PRIOR_AORSURG				
Both	_	_		
Endo	1.36	$0.52,\ 3.53$	0.53	
None	1.04	0.41, 2.61	0.94	
Open	0.99	0.36, 2.76	0.99	
PRIOR_CHF				
No	_	_		
Yes	1.08	0.76, 1.54	0.66	
PREOP_GFR_CAT				
End-stage renal disease	_	_		
Mildly decreased	1.83	0.23, 14.4	0.56	
Mildly to severely decreased	2.51	0.32, 19.7	0.38	
Normal or increased	2.12	0.27, 16.9	0.48	
Severely decreased	2.08	0.24,18.3	0.51	
TREATED_RENALS				
No				
Yes	0.54	0.39, 0.73	< 0.001	
PATHOLOGY				
Aneurysm	_	_		
Aneurysm from dissection	0.33	0.10, 1.09	0.068	
Dissection	0.85	0.31, 2.33	0.76	
PAU/IMH	1.21	0.35, 4.23	0.77	
extent				
Juxtarenal	_	_		
No	1.12	0.82, 1.54	0.47	

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

 ${\bf POSTOP_INTISCH:\ Post-op\ Intestinal\ Ischemia}$

	Intestinal Ischemia				
Characteristic	OR^1	95% CI ¹	p-value		
PRESENTATION					
Asymptomatic		_			
Symptomatic	1.13	0.56, 2.26	0.74		

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

	Intestinal Ischemia			
Characteristic	OR^1	95% CI ¹	p-value	
PRESENTATION	'			
Asymptomatic	_	_		
Symptomatic	1.07	0.50, 2.30	0.87	
AGECAT				
< 50	_	_		
>79	0.41	0.07, 2.37	0.32	
50-59	0.49	0.06, 4.15	0.51	
60-69	0.36	0.06, 2.15	0.26	
70-79	0.45	0.08, 2.48	0.36	
GENDER				
female	_	_		
male	0.62	0.37, 1.03	0.063	
PREOP_SMOKING				
No				
Yes	0.86	0.43, 1.70	0.66	
PRIOR_AORSURG				
Both	_			
Endo	1.94	0.23, 16.6	0.55	
None	1.84	0.22,15.8	0.58	
Open	1.62	0.16, 15.9	0.68	
PRIOR_CHF				
No	_	_		
Yes	1.05	0.53, 2.08	0.89	
PREOP_DIALYSIS				

 $^{^{1}\}mathrm{OR}=\mathrm{Odds}$ Ratio, CI = Confidence Interval

	Intestinal Ischemia			
Characteristic	$order R^1$	95% CI ¹	p-value	
No	_	_		
Yes	0.69	0.09, 5.12	0.71	
PATHOLOGY				
Aneurysm	_	_		
Aneurysm from dissection	0.48	0.06, 3.66	0.48	
Dissection	0.71	$0.14,\ 3.71$	0.69	
PAU/IMH	1.28	0.16, 10.6	0.82	
extent				
Juxtarenal	_	_		
No	0.94	0.53, 1.69	0.85	

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

POSTOP_CEREBROSX: Post-op Cerebrovascular Stroke:

Also account for ARMNECK_ACCESS

	Post-op Cerebrovascular Stroke				
Characteristic	OR^1	95% CI 1	p-value		
PRESENTATION					
Asymptomatic		_			
Symptomatic	3.42	1.83,6.38	< 0.001		

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

		Cerebrovaso	cular Stroke
Characteristic	\mathbf{OR}^1	$95\%~\mathrm{CI}^1$	p-value
PRESENTATION			
Asymptomatic		_	
Symptomatic	1.81	0.91,3.60	0.091
AGECAT			
< 50		_	
>79	0.51	0.08, 3.29	0.48
50-59	0.79	0.13,4.95	0.81
60-69	0.23	0.04,1.45	0.12
70-79	0.62	0.11, 3.46	0.58
GENDER			
female	_	_	
male	0.49	0.26, 0.90	0.023
PREOP_SMOKING			
No		_	
Yes	0.73	0.34, 1.58	0.43
PRIOR_AORSURG			
Both	_	_	
Endo	1.20	0.24,6.05	0.83
None	1.39	0.28, 6.92	0.69
Open	0.95	0.15, 5.88	0.95
PRIOR_CHF			
No		_	
Yes	2.14	1.07, 4.31	0.032
$^{1}OR = Odds Ratio, CI = Conf$	idence Inter	rval	

	Post-op	Cerebrovaso	cular Stroke
Characteristic	OR^1	95% CI 1	p-value
PREOP_DIALYSIS			
No		_	
Yes	1.97	0.46, 8.55	0.36
PATHOLOGY			
Aneurysm	_	_	
Aneurysm from dissection	3.44	1.03, 11.6	0.045
Dissection	2.56	0.57, 11.5	0.22
PAU/IMH	1.46	$0.21,\ 10.2$	0.70
extent			
Juxtarenal		_	
No	1.44	0.68, 3.04	0.34
ARMNECK_ACCESS			
No	_	_	
Yes	2.31	1.26, 4.25	0.007

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

POSTOP_SPINAL_ISCHEMIA: Post-op Spinal Ischemia

Also account for ${\tt POSTOP_SPINALDRAIN}$ and ${\tt OCCLUDED_CELIAC}$

	Post-op Spinal Ischemia				
Characteristic	OR^1	95% CI 1	p-value		
PRESENTATION	,				
Asymptomatic					
Symptomatic	2.56	1.70, 3.86	< 0.001		

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

-			
	Post-op Spinal Ischen		
Characteristic	\mathbf{OR}^1	$95\%~\mathrm{CI}^1$	p-value
PRESENTATION	,		
Asymptomatic		_	
Symptomatic	1.61	1.00, 2.60	0.050
AGECAT			
< 50	_	_	
>79	2.19	$0.27,\ 17.7$	0.46
50-59	1.99	0.23,17.3	0.53
60-69	1.63	0.21,13.0	0.64
70-79	2.13	0.27, 16.6	0.47
GENDER			
female	_	_	
male	0.88	0.58,1.36	0.58
PREOP_SMOKING			
No			
Yes	1.22	0.64, 2.32	0.55
PRIOR_AORSURG			
Both	_	_	
Endo	1.23	0.52,2.95	0.64
None	1.46	0.63, 3.40	0.37
Open	1.47	0.60, 3.63	0.40
PRIOR_CHF			
No			
Yes	1.06	0.61, 1.83	0.84
$^{1}OR = Odds Ratio, CI = Confid$	dence Inter	rval	

	Pos	t-op Spinal I	schemia
Characteristic	OR^1	95% CI ¹	p-value
PREOP_DIALYSIS			
No		_	
Yes	1.16	0.48, 2.83	0.75
PATHOLOGY			
Aneurysm		_	
Aneurysm from dissection	0.98	0.43, 2.25	0.96
Dissection	1.08	0.36,3.26	0.89
PAU/IMH	0.44	0.05, 3.63	0.45
extent			
Juxtarenal		_	
No	1.01	0.65, 1.57	0.97
POSTOP_SPINALDRAIN			
No		_	
Yes	19.0	11.1, 32.6	< 0.001
OCCLUDED_CELIAC			
FALSE	_	_	
TRUE	1.57	0.83, 2.99	0.17

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

POSTOP_LOS: Length of stay in days between surgery date and discharge date

	Length of stay in days between surgery date and discharge date				
Characteristic	OR^1	$95\%~\mathrm{CI}^1$	p-value		
PRESENTATION					
Asymptomatic		_			
Symptomatic	3.39	2.69, 4.29	< 0.001		

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

		of stay in da date and dis	
Characteristic	OR^1	95% CI ¹	p-value
PRESENTATION			
Asymptomatic	_	_	
Symptomatic	2.46	$1.91,\ 3.17$	< 0.001
AGECAT			
< 50			
>79	2.28	0.88, 5.88	0.089
50-59	1.93	0.72, 5.21	0.19
60-69	1.55	0.60, 3.96	0.36
70-79	2.49	0.98,6.35	0.055
GENDER			
female		_	
male	0.62	0.50, 0.75	< 0.001
PREOP_SMOKING			
No	_	_	
Yes	0.97	0.74, 1.28	0.82
PRIOR_AORSURG			
Both		_	
Endo	1.26	0.73, 2.19	0.41
None	0.96	0.57, 1.62	0.88
Open	1.36	0.76, 2.41	0.30
PRIOR_CHF			
No	_	_	
Yes	1.41	1.11, 1.80	0.005
1 OR = Odds Ratio, CI = Confi	dence Inter	val	

¹⁵

	Length of stay in days between surgery date and discharge dat		
Characteristic	OR^1	$95\%~\mathrm{CI}^1$	p-value
PREOP_DIALYSIS			
No	_	_	
Yes	1.74	1.00, 3.00	0.048
PATHOLOGY			
Aneurysm		_	
Aneurysm from dissection	1.59	0.96, 2.63	0.069
Dissection	1.96	$1.11,\ 3.45$	0.020
PAU/IMH	1.82	0.90, 3.67	0.094
extent			
Juxtarenal			
No	1.97	1.60, 2.44	< 0.001

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

 ${\bf POSTOP_COMPLICATIONS:\ Any\ Complications\ Post-op}$

	Any Complications Post-op			
Characteristic	OR^1	95% CI ¹	p-value	
PRESENTATION				
Asymptomatic		_		
Symptomatic	1.74	1.37, 2.21	< 0.001	

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

	Any Complications Post-			
Characteristic	OR^1	95% CI ¹	p-value	
PRESENTATION	,			
Asymptomatic	_	_		
Symptomatic	1.44	1.12, 1.86	0.004	
AGECAT				
< 50	_	_		
>79	1.17	0.46, 2.97	0.74	
50-59	1.29	0.49, 3.42	0.61	
60-69	0.97	0.39, 2.41	0.94	
70-79	1.32	0.53, 3.31	0.55	
GENDER				
female	_	_		
male	0.65	0.54, 0.78	< 0.001	
PREOP_SMOKING				
No				
Yes	1.01	0.79, 1.31	0.91	
PRIOR_AORSURG				
Both				
Endo	0.95	0.56, 1.60	0.84	
None	1.00	0.60, 1.65	0.99	
Open	1.41	0.82, 2.42	0.21	
PRIOR_CHF				
No				
Yes	1.28	1.04, 1.58	0.018	
PREOP_DIALYSIS				

 $^{^{1}\}mathrm{OR}=\mathrm{Odds}$ Ratio, CI = Confidence Interval

	Any (Complication	s Post-op
Characteristic	OR^1	95% CI ¹	p-value
No	_	_	,
Yes	1.25	0.73, 2.15	0.41
PATHOLOGY			
Aneurysm		_	
Aneurysm from dissection	1.32	0.82,2.12	0.26
Dissection	0.94	0.51,1.73	0.84
PAU/IMH	0.77	0.35, 1.70	0.52
extent			
Juxtarenal	_	_	
No	1.47	$1.21,\ 1.78$	< 0.001

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

 $RETX_R_RTOR \hbox{: } Re\hbox{-intervention}$

	Re-intervention				
Characteristic	OR^1	95% CI ¹	p-value		
PRESENTATION					
Asymptomatic		_			
Symptomatic	1.86	1.33, 2.62	< 0.001		

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

	,	Re-intervent	tion
Characteristic	OR^1	95% CI 1	p-value
PRESENTATION			
Asymptomatic	_	_	
Symptomatic	1.48	1.04, 2.09	0.029
AGECAT			
< 50	_	_	
>79	0.95	$0.28,\ 3.27$	0.93
50-59	1.41	0.38, 5.19	0.60
60-69	0.70	0.20, 2.44	0.58
70-79	0.94	0.28, 3.14	0.92
GENDER			
female			
male	0.59	0.45, 0.78	< 0.001
PREOP_SMOKING			
No	_	_	
Yes	1.29	0.81, 2.05	0.28
PRIOR_AORSURG			
Both	_	_	
Endo	0.95	0.43, 2.09	0.90
None	1.01	0.47, 2.16	0.99
Open	0.67	0.27, 1.63	0.37
PRIOR_CHF			
No	_	_	
Yes	1.21	0.86, 1.72	0.27
PREOP_DIALYSIS			

 $^{^{1}\}mathrm{OR}=\mathrm{Odds}$ Ratio, $\mathrm{CI}=\mathrm{Confidence}$ Interval

	'	Re-intervent	tion
Characteristic	OR^1	95% CI ¹	p-value
No	_	_	
Yes	0.75	0.31, 1.81	0.52
PATHOLOGY			
Aneurysm	_	_	
Aneurysm from dissection	1.68	0.88, 3.19	0.12
Dissection	0.64	0.23,1.78	0.39
PAU/IMH	1.77	0.70, 4.47	0.23
extent			
Juxtarenal	_	_	
No	1.56	1.15, 2.12	0.004

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

BRANCH_POST: Post-treatment Status of All Branches

	Post-treatment Status of All Branches			
Characteristic	OR^1	95% CI ¹	p-value	
PRESENTATION				
Asymptomatic				
Symptomatic	2.28	1.78, 2.91	< 0.001	

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

	Post-treatment Status of All Branches			
Characteristic	OR^1	95% CI ¹	p-value	
PRESENTATION				
Asymptomatic	_	_		
Symptomatic	1.64	1.24, 2.16	< 0.001	
AGECAT				
< 50	_	_		
>79	0.78	0.32, 1.90	0.58	
50-59	1.00	0.39, 2.58	>0.99	
60-69	0.95	0.40, 2.27	0.91	
70-79	0.90	0.38, 2.14	0.81	
GENDER				
female				
male	0.47	0.39, 0.58	< 0.001	
PREOP_SMOKING				
No				
Yes	1.08	0.79, 1.48	0.63	
PRIOR_AORSURG				
Both	_	_		
Endo	0.79	0.46, 1.35	0.38	
None	0.52	0.31, 0.86	0.011	
Open	0.69	0.39, 1.23	0.21	
PRIOR_CHF				
No	_	_		
Yes	1.43	1.12, 1.82	0.004	
¹ OR = Odds Ratio, CI = Confidence Interval				

 $^{{}^{1}}OR = Odds Ratio, CI = Confidence Interval$

	Post-treatment Status of All Branches			
Characteristic	OR^1	95% CI ¹	p-value	
PREOP_DIALYSIS				
No	_	_		
Yes	7.90	4.70, 13.3	< 0.001	
PATHOLOGY				
Aneurysm	_	_		
Aneurysm from dissection	0.77	0.43,1.37	0.37	
Dissection	1.57	0.90, 2.74	0.12	
PAU/IMH	0.45	0.16, 1.24	0.12	
extent				
Juxtarenal	_	_		
No	1.56	1.25, 1.93	< 0.001	

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

For the following variables, we need to merge age group "<50" and "50-59" so that there's no 0 patients on one level.

POSTOP_DIALYSIS: Post-op Dialysis

Also account for OCCLUDED_RENAL

	Post-op Dialysis			
Characteristic	OR^1	$95\%~\mathrm{CI}^1$	p-value	
PRESENTATION	,		,	
Asymptomatic		_		
Symptomatic	2.42	1.44,4.07	< 0.001	

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

	Post-op Dialysis			
Characteristic	$ormalfont{OR}^1$	$95\%~\mathrm{CI}^1$	p-value	
PRESENTATION				
Asymptomatic	_	_		
Symptomatic	1.78	1.01, 3.13	0.047	
AGECAT				
< 59		_		
60-69	0.53	0.22,1.27	0.16	
70-79	0.40	0.17, 0.97	0.042	
>79	0.39	0.15, 1.01	0.053	
GENDER				
female				
male	0.62	0.38, 1.00	0.049	
PREOP_SMOKING				
No				
Yes	1.31	0.60, 2.86	0.49	
PRIOR_AORSURG				
Both		_		
Endo	1.04	0.26, 4.10	0.95	
None	0.82	0.22, 3.04	0.76	
Open	0.58	0.12, 2.80	0.50	
PRIOR_CHF				
No				
10D 011 D // OT 0 C1	T .	1		

 $^{^{1}\}mathrm{OR}=\mathrm{Odds}$ Ratio, CI = Confidence Interval

	Post-op Dialysis			
Characteristic	OR^1	95% CI ¹	p-value	
Yes	0.74	0.37, 1.45	0.38	
PREOP_DIALYSIS				
No	_	_		
Yes	0.00	0.00, 0.00	< 0.001	
PATHOLOGY				
Aneurysm	_	_		
Aneurysm from dissection	0.33	0.04, 2.96	0.32	
Dissection	1.74	0.52, 5.82	0.37	
PAU/IMH	2.33	0.51, 10.6	0.27	
extent				
Juxtarenal	_	_		
No	0.99	0.58, 1.70	0.98	
OCCLUDED_RENAL				
FALSE	_	_		
TRUE	4.82	2.33, 9.95	< 0.001	

 $[\]overline{^{1}\text{OR} = \text{Odds Ratio, CI} = \text{Confidence Interval}}$

 ${\bf POSTOP_LEGEMBO: Leg~Ischemia/Embol}$

	Leg Ischemia/Embol			
Characteristic	OR^1	95% CI ¹	p-value	
PRESENTATION				
Asymptomatic		_		
Symptomatic	1.97	1.13, 3.46	0.017	

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

	Leg Ischemia/Embol			
Characteristic	OR^1	95% CI ¹	p-value	
PRESENTATION				
Asymptomatic	_	_		
Symptomatic	1.74	0.94, 3.22	0.078	
AGECAT				
< 59	_			
60-69	0.80	0.28, 2.27	0.68	
70-79	0.80	0.29, 2.23	0.67	
>79	0.67	0.23, 2.00	0.48	
GENDER				
female	_	_		
male	0.43	0.28, 0.67	< 0.001	
PREOP_SMOKING				
No	_	_		
Yes	2.61	0.98, 6.96	0.056	
PRIOR_AORSURG				
Both	_	_		
Endo	139,944,960,652,840			
None	$162,\!086,\!767,\!451,\!715$			
Open	288,216,346,513,340			
PRIOR_CHF				
No	_			
Yes	0.85	0.44, 1.64	0.63	
PREOP_DIALYSIS				
No	_	_		
1 OR = Odds Ratio, CI = Co	nfidence Interval			

	Leg Is		
Characteristic	OR^1	$95\%~\mathrm{CI}^1$	p-value
Yes	0.58	0.08, 4.19	0.59
PATHOLOGY			
Aneurysm	_	_	
Aneurysm from dissection	0.87	0.20, 3.86	0.86
Dissection	0.60	0.08, 4.59	0.62
PAU/IMH	2.28	$0.49,\ 10.6$	0.29
extent			
Juxtarenal	_	_	
No	1.01	0.61, 1.66	0.98

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

 ${\bf POSTOP_RESPIRATORY:\ Post-op\ Respiratory}$

	Post-op Respiratory			
Characteristic	OR^1	95% CI 1	p-value	
PRESENTATION				
Asymptomatic				
Symptomatic	2.65	1.87, 3.75	< 0.001	

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

	Post-op Respiratory			
Characteristic	$ormalfont{OR}^1$	$95\%~\mathrm{CI}^1$	p-value	
PRESENTATION				
Asymptomatic				
Symptomatic	2.17	1.47, 3.21	< 0.001	
AGECAT				
<59				
60-69	0.94	0.44,1.98	0.86	
70-79	1.06	0.50, 2.25	0.87	
>79	0.94	0.42, 2.06	0.87	
GENDER				
female	_	_		
male	0.66	0.47, 0.92	0.014	
PREOP_SMOKING				
No	_	_		
Yes	1.08	0.68, 1.72	0.75	
PRIOR_AORSURG				
Both	_	_		
Endo	0.59	0.27, 1.30	0.19	
None	0.55	0.26, 1.16	0.11	
Open	0.67	0.28, 1.60	0.36	
PRIOR_CHF				
No	_	_		
Yes	1.23	0.82,1.84	0.31	
PREOP_DIALYSIS				
No	_			
$^{1}\mathrm{OR}=\mathrm{Odds}\;\mathrm{Ratio},\mathrm{CI}=\mathrm{Con}$	fidence Inter	val		

	Post-op Respiratory		
Characteristic	OR^1	95% CI ¹	p-value
Yes	0.78	0.27, 2.23	0.64
PATHOLOGY			
Aneurysm			
Aneurysm from dissection	0.60	0.23,1.55	0.29
Dissection	0.83	0.29, 2.38	0.73
PAU/IMH	1.14	0.33, 4.02	0.83
extent			
Juxtarenal	_	_	
No	1.74	1.25, 2.41	< 0.001

 $[\]overline{^{1}}$ OR = Odds Ratio, CI = Confidence Interval

POSTOP_PRBC: Transfusion # Units PRBC

We don't include this variable in the final report.

Code Appendix

```
knitr::opts_chunk$set(echo = FALSE,message = FALSE,warning = FALSE)
library(tidyverse)
library(table1)
library(dplyr)
library(geepack)
library(gtsummary)
library(broom.mixed)
## ----- working directories for Lily -----
wd_lily = '/Users/hanyiwang/Desktop/Comparative-analysis-of-treatments-of-CAA'
path_lily = c("../data/TEVAR_PROC.csv")
## ----- working directories for Jenn ------
wd_jenn = '/Users/jenniferci/Desktop/stlp new laptop/Capstone/Comparative-analysis-of-treatments-of-CAA
path_jenn = c("TEVAR_PROC.csv")
## ----- read data -----
setwd(wd_lily)
TEVAR_PROC = read.csv(path_lily)
# setwd(wd_jenn)
# PROC = read.csv(path_jenn)
# ### ----- Correlation matrix -----
# library(corrplot)
# library(tidyverse)
# library(caret)
# library(ggcorrplot)
#
# matrix <- PROC %>%
     select_if(is.numeric) %>% subset(., select = -1)%>%
     cor(.,use = "complete")
#
#
# corrplot(matrix, method="number")
# #select dataset that column are not numeric
# matrix <- PROC %>% select if(negate(is.numeric))
# # find out the variables
# lapply(matrix[,], unique)
# sapply(lapply(matrix, unique), length)
# #select if more than one variable
# y<-matrix %>%select_if(function(col) length(unique(col))==2)
# # Convert all columns to factor
# data3 <- as.data.frame(unclass(y),</pre>
                        stringsAsFactors = TRUE)
#
# model.matrix(~0 +., data=data3) %>%
#
  cor(use="pairwise.complete.obs") %>%
#
   ggcorrplot(show.diag = F, type="lower", lab=TRUE, lab_size=2)
```

```
# str(y)
## ----- modify dataset -----
TEVAR_PROC = TEVAR_PROC %>%
  mutate(extent = factor(extent,levels = c("Juxtarenal AAA", "Type 1 TAAA", "Type 2 TAAA", "Type 3 TAAA", "
  mutate(GFRCHANGE = (PREOP_GFR-POSTOP_GFR)/PREOP_GFR) %>%
  mutate(GFRFIFTY = (ifelse(GFRCHANGE<0.5, "<50%",</pre>
                            ifelse(GFRCHANGE>=0.5, ">=50%",NA)))) %>%
  mutate(TREATED_RENALS = (ifelse(NUM_TREATED_RENALS<1, "No",</pre>
                                  ifelse(NUM_TREATED_RENALS>=1, "Yes",NA)))) %>%
  mutate(TOTAL_LOS = case_when(TOTAL_LOS>7 ~ '>7',
                               TOTAL_LOS<=7 ~ '<=7')) %>%
  mutate(ICUSTAY = case_when(ICUSTAY>4 ~ '>4',
                             ICUSTAY<=4 ~ '<=4')) %>%
  mutate(POSTOP_LOS = case_when(POSTOP_LOS>7 ~ '>7',
                                POSTOP_LOS<=7 ~ '<=7')) %>%
  mutate(TOTAL_LOS = as.numeric(TOTAL_LOS != "<=7") ) %>%
  mutate(ICUSTAY = as.numeric(ICUSTAY != "<=4") ) %>%
  mutate(POSTOP_LOS = as.numeric(POSTOP_LOS != "<=7") ) %>%
  mutate(POSTOP_INTISCH = as.numeric(POSTOP_INTISCH == "Yes")) %>%
  mutate(POSTOP CEREBROSX = as.numeric(POSTOP CEREBROSX == "Yes")) %>%
  mutate(POSTOP_SPINAL_ISCHEMIA = as.numeric(POSTOP_SPINAL_ISCHEMIA == "Yes")) %>%
  mutate(POSTOP_DIALYSIS = as.numeric(POSTOP_DIALYSIS == "Yes")) %>%
  mutate(POSTOP_COMPLICATIONS = as.numeric(POSTOP_COMPLICATIONS == "Yes")) %>%
  mutate(POSTOP_LEGEMBO = as.numeric(POSTOP_LEGEMBO == "Yes")) %>%
  mutate(POSTOP_RESPIRATORY = as.numeric(POSTOP_RESPIRATORY == "Yes")) %>%
  mutate(RETX_R_RTOR = as.numeric(RETX_R_RTOR == "Yes")) %>%
  mutate(BRANCH_POST = as.numeric(BRANCH_POST == "Yes")) %>%
  mutate(GFRFIFTY = as.numeric(GFRFIFTY != "<50%") )</pre>
# remove missing values
TEVAR_PROC = subset(TEVAR_PROC, !is.na(extent))
## ----- change class of variables -----
names <- c('CENTERID','AGECAT', 'GENDER', 'PREOP_SMOKING', 'PRIOR_AORSURG', 'PRIOR_CHF', 'PREOP_DIALYSI</pre>
TEVAR_PROC[,names] <- lapply(TEVAR_PROC[,names] , factor)</pre>
## ------ Length of stay in days between admission date and discharge date -----
#table(PROC$extent)
#PROCNM[is.na(PROCNM) | PROCNM=="Inf"] = NA
# repeated patients, need id, geeglm needs complete data, extent is not complete
## ----- unadjusted -----
LOS1<-geeglm(TOTAL_LOS~ PRESENTATION, data=TEVAR_PROC, family=binomial(link="logit"), id = CENTERID, co.
t1<-LOS1 %>%
 tbl_regression(exponentiate=TRUE,
```

```
tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**TOTAL_LOS Length of Stay in days**")%>%as_flex_table()
## ----- adjusted -----
LOS2<-geeglm(TOTAL_LOS ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+PRIOR_AORSURG+PRIOR_CHF+PREOP_DIALYSI
            , data=TEVAR_PROC, family=binomial(link="logit"),id = CENTERID, corstr = "independence")
t2<-LOS2 %>%
 tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold_p(t = 0.05)
tbl_merge(tbls = list(t2),tab_spanner ="**TOTAL_LOS Length of Stay in days**")%>%as_flex_table()
#the difference in mean number of sessions attended comparing treatment to control
#adjust<-c('PRESENTATION','AGECAT', 'GENDER', 'PREOP_SMOKING','PRIOR_AORSURG', 'PRIOR_CHF','PATHOLOGY',
#PROC %>%
# tbl_uvregression(
    y = TOTAL\_LOS,
    x = PRESENTATION+AGECAT+GENDER+PREOP SMOKING+PRIOR AORSURG+PRIOR CHF+PATHOLOGY+NUM TREATED BRANCHE
   method = qeepack::qeeqlm,
   method.args = list(id = X+CENTERID, corstr = "independence"),
    include = all_of(adjust)
# ) %>%
# as_kable()
## ----- ICU Stay -----
## ----- unadjusted -----
ICU1<-geeglm(ICUSTAY~ PRESENTATION, data=TEVAR_PROC, family=binomial(link="logit"), id = CENTERID, cors
#summary(LOS2)
#summary(ICU1)
#ICU1<-geeglm(ICUSTAY ~ PRESENTATION, data=PROCNM, id = CENTERID, corstr = "independence")
t1<-ICU1 %%tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                          pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold_p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**ICU Stay**")%>%as_flex_table()
## ----- adjusted -----
ICU2<-geeglm(ICUSTAY ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+PRIOR_AORSURG+PRIOR_CHF+PREOP_DIALYSIS+
            , family=binomial(link="logit"), data=TEVAR_PROC, id = CENTERID, corstr = "independence")
t2<- ICU2 %>%tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                           pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold_p(t = 0.05)
```

```
tbl_merge(tbls = list(t2),tab_spanner ="**ICU Stay**")%>%as_flex_table()
## ----- Creatinine -----
## ----- unadjusted -----
POSTOP_HIGHCREAT1<-geeglm(POSTOP_HIGHCREAT ~ PRESENTATION, data=TEVAR_PROC, id = CENTERID, corstr = "ind
t1<-POSTOP HIGHCREAT1 %>%
 tbl_regression(tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**Highest Creatinine**")%>%as_flex_table()
## ----- adjusted -----
POSTOP_HIGHCREAT2<-geeglm(POSTOP_HIGHCREAT ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+PRIOR_AORSURG+PRI
                        , data=TEVAR_PROC, id = CENTERID, corstr = "independence")
t2<- POSTOP_HIGHCREAT2 %>%
 tbl_regression(tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold_p(t = 0.05)
tbl_merge(tbls = list(t2),tab_spanner ="**Highest Creatinine**")%>%as_flex_table()
## ----- GFR reduction -----
#table(PROC$GFRFIFTY)
## ----- unadjusted -----
GFR1<-geeglm(GFRFIFTY~ PRESENTATION, data=TEVAR_PROC, family=binomial(link="logit"), id = CENTERID, cor
t1<-GFR1 %>%
 tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                         pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold_p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**GFR reduction**")%>%as_flex_table()
## ----- adjusted -----
GFR2<-geeglm(GFRFIFTY ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+PRIOR_AORSURG+PRIOR_CHF+PREOP_GFR_CAT+
           , family=binomial(link="logit"), data=TEVAR_PROC, id = CENTERID, corstr = "independence")
 tbl_regression(exponentiate=TRUE,tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold_p(t = 0.05)
tbl_merge(tbls = list(t2),tab_spanner ="**GFR reduction**")%>%as_flex_table()
## ----- Post-op Intestinal Ischemia -----
## ----- unadjusted -----
POSTOP_INTISCH1<-geeglm(POSTOP_INTISCH ~ PRESENTATION, family=binomial(link="logit"), data=TEVAR_PROC,i
t1<-POSTOP_INTISCH1 %>%tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                                    pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold_p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**Intestinal Ischemia**")%>%as_flex_table()
```

```
## ----- adjusted -----
POSTOP_INTISCH2<-geeglm(POSTOP_INTISCH ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+PRIOR_AORSURG+PRIOR_C
                                           , data=TEVAR_PROC, family=binomial(link="logit"), id = CENTERID, corstr = "independent of the content of the corst of the 
t2<- POSTOP_INTISCH2 %>%tbl_regression(exponentiate=TRUE, tidy_fum = broom.mixed::tidy,
                                                                        pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
    bold_p(t = 0.05)
tbl_merge(tbls = list(t2),tab_spanner ="**Intestinal Ischemia**")%>%as_flex_table()
# model<-geeglm(as.numeric(POSTOP_DIALYSIS)~PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+PRIOR_AORSURG+PRIO
# model <- queqlm (outcome ~ predictor + confounder, family = binomial (link = "logit"),
# data=na.omit(data), corstr='ar1', id=id, std.err="san.se")
# TOTAL_LOS
# ICUSTAY
# POSTOP_PRBC
# POSTOP_HIGHCREAT
# POSTOP_INTISCH
## ----- Post-op Cerebrovascular Stroke -----
## ----- unadjusted -----
POSTOP_CEREBROSX1 = geeglm(POSTOP_CEREBROSX ~ PRESENTATION,
                                                  data=TEVAR_PROC, family=binomial(link="logit"),
                                                  id = CENTERID, corstr = "independence")
t1 = POSTOP_CEREBROSX1 %>%
   tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
    bold_p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**Post-op Cerebrovascular Stroke**")%>%as_flex_table()
## ----- adjusted -----
POSTOP_CEREBROSX2 = geeglm(POSTOP_CEREBROSX ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+
                                        PRIOR_AORSURG+PRIOR_CHF+PREOP_DIALYSIS+PATHOLOGY+extent+ARMNECK_ACCESS,
                                     family=binomial(link="logit"), data=TEVAR_PROC,
                                     id = CENTERID, corstr = "independence")
t2 = POSTOP CEREBROSX2 %>%
   tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
    bold_p(t = 0.05)
tbl_merge(tbls = list(t2),tab_spanner ="**Post-op Cerebrovascular Stroke**")%>%as_flex_table()
## ----- Post-op Spinal Ischemia -----
## ----- unadjusted -----
POSTOP_SPINAL_ISCHEMIA1 = geeglm(POSTOP_SPINAL_ISCHEMIA ~ PRESENTATION,
                                                             data=TEVAR_PROC,family=binomial(link="logit"),
                                                             id = CENTERID, corstr = "independence")
t1 = POSTOP_SPINAL_ISCHEMIA1 %>%
    tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
    bold_p(t = 0.05)
```

```
tbl_merge(tbls = list(t1),tab_spanner ="**Post-op Spinal Ischemia**")%>%as_flex_table()
## ----- adjusted -----
POSTOP SPINAL ISCHEMIA2 = geeglm(POSTOP SPINAL ISCHEMIA ~ PRESENTATION+AGECAT+GENDER+
                                  PREOP_SMOKING+PRIOR_AORSURG+PRIOR_CHF+
                                  PREOP DIALYSIS+PATHOLOGY+extent+
                                  POSTOP_SPINALDRAIN+OCCLUDED_CELIAC,
                   family=binomial(link="logit"), data=TEVAR_PROC,
                   id = CENTERID, corstr = "independence")
t2 = POSTOP_SPINAL_ISCHEMIA2 %>%
  tbl_regression(exponentiate=TRUE,tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
  bold_p(t = 0.05)
tbl_merge(tbls = list(t2),tab_spanner ="**Post-op Spinal Ischemia**")%>%as_flex_table()
## ------ Length of stay in days between surgery date and discharge date ------
## ----- unadjusted -----
POSTOP_LOS1 = geeglm(POSTOP_LOS~ PRESENTATION,
                    data=TEVAR_PROC, family=binomial(link="logit"),
                    id = CENTERID, corstr = "independence")
# summary(POSTOP_LOS1)
t1 = POSTOP_LOS1 %>%
 tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
  bold_p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**Length of stay in days between surgery date and discharge dat
## ----- adjusted -----
POSTOP_LOS2 = geeglm(POSTOP_LOS ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+
                     PRIOR_AORSURG+PRIOR_CHF+PREOP_DIALYSIS+PATHOLOGY+extent,
                   family=binomial(link="logit"), data=TEVAR_PROC,
                   id = CENTERID, corstr = "independence")
t2 = POSTOP LOS2 %>%
 tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold_p(t = 0.05)
tbl_merge(tbls = list(t2),tab_spanner ="**Length of stay in days between surgery date and discharge dat
## ----- Any Complications Post-op -----
## ----- unadjusted -----
POSTOP_COMPLICATIONS1 = geeglm(POSTOP_COMPLICATIONS~ PRESENTATION,
                    data=TEVAR_PROC, family=binomial(link="logit"),
                    id = CENTERID, corstr = "independence")
t1 = POSTOP_COMPLICATIONS1 %>%
 tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
```

```
pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold_p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**Any Complications Post-op**")%>%as_flex_table()
## ----- adjusted -----
POSTOP_COMPLICATIONS2 = geeglm(POSTOP_COMPLICATIONS ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+
                     PRIOR AORSURG+PRIOR CHF+PREOP DIALYSIS+PATHOLOGY+extent,
                   family=binomial(link="logit"), data=TEVAR_PROC,
                   id = CENTERID, corstr = "independence")
t2 = POSTOP_COMPLICATIONS2 %>%
 tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold_p(t = 0.05)
tbl_merge(tbls = list(t2),tab_spanner ="**Any Complications Post-op**")%>%as_flex_table()
## ----- Re-intervention -----
## ----- unadjusted -----
RETX_R_RTOR1 = geeglm(RETX_R_RTOR~ PRESENTATION,
                    data=TEVAR_PROC, family=binomial(link="logit"),
                    id = CENTERID, corstr = "independence")
# summary(POSTOP_LOS1)
t1 = RETX R RTOR1 %>%
 tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold_p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**Re-intervention**")%>%as_flex_table()
## ----- adjusted -----
RETX_R_RTOR2 = geeglm(RETX_R_RTOR ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+
                     PRIOR_AORSURG+PRIOR_CHF+PREOP_DIALYSIS+PATHOLOGY+extent,
                   family=binomial(link="logit"), data=TEVAR_PROC,
                   id = CENTERID, corstr = "independence")
t2 = RETX_R_RTOR2 %>%
 tbl_regression(exponentiate=TRUE,tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
 bold_p(t = 0.05)
tbl merge(tbls = list(t2), tab spanner ="**Re-intervention**")%>%as flex table()
## ----- Post-treatment Status of All Branches ------
## ----- unadjusted -----
BRANCH_POST1 = geeglm(BRANCH_POST~ PRESENTATION,
                    data=TEVAR_PROC, family=binomial(link="logit"),
                    id = CENTERID, corstr = "independence")
# summary(POSTOP_LOS1)
t1 = BRANCH_POST1 %>%
 tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
```

```
bold_p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**Post-treatment Status of All Branches**")%>%as_flex_table()
## ----- adjusted -----
BRANCH_POST2 = geeglm(BRANCH_POST ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+
                     PRIOR_AORSURG+PRIOR_CHF+PREOP_DIALYSIS+PATHOLOGY+extent,
                   family=binomial(link="logit"), data=TEVAR PROC,
                   id = CENTERID, corstr = "independence")
t2 = BRANCH_POST2 %>%
  tbl_regression(exponentiate=TRUE,tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
  bold_p(t = 0.05)
tbl_merge(tbls = list(t2),tab_spanner ="**Post-treatment Status of All Branches**")%>%as_flex_table()
## ----- modify dataset -----
TEVAR_PROC = TEVAR_PROC %>%
  mutate(AGECAT = factor(AGECAT,levels = c('<50','50-59','60-69','70-79','>79'),
                        labels = c('<59','<59','60-69','70-79','>79')))
## ----- Post-op Dialysis -----
# table1(~ AGECAT | factor(POSTOP_DIALYSIS),data=subset(TEVAR_PROC, !is.na(POSTOP_DIALYSIS)))
## ----- unadjusted -----
POSTOP_DIALYSIS1 = geeglm(POSTOP_DIALYSIS ~ PRESENTATION,
                                data=TEVAR_PROC,family=binomial(link="logit"),
                                id = CENTERID, corstr = "independence")
t1 = POSTOP_DIALYSIS1 %>%
  tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
  bold_p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**Post-op Dialysis**")%>%as_flex_table()
## ----- adjusted -----
POSTOP_DIALYSIS2 = geeglm(POSTOP_DIALYSIS ~ PRESENTATION+AGECAT+GENDER+
                                  PREOP_SMOKING+PRIOR_AORSURG+PRIOR_CHF+
                                  PREOP_DIALYSIS+PATHOLOGY+extent+
                                  OCCLUDED_RENAL,
                   family=binomial(link="logit"), data=TEVAR_PROC,
                   id = CENTERID, corstr = "independence")
t2 = POSTOP_DIALYSIS2 %>%
 tbl_regression(exponentiate=TRUE,tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
  bold_p(t = 0.05)
tbl_merge(tbls = list(t2),tab_spanner ="**Post-op Dialysis**")%>%as_flex_table()
## ----- Leg Ischemia/Emboli -----
# table1(~ AGECAT | factor(POSTOP_LEGEMBO),data=subset(TEVAR_PROC, !is.na(POSTOP_LEGEMBO)))
```

```
## ----- unadjusted -----
POSTOP_LEGEMB01 = geeglm(POSTOP_LEGEMB0~ PRESENTATION,
                    data=TEVAR PROC, family=binomial(link="logit"),
                    id = CENTERID, corstr = "independence")
# summary(POSTOP LOS1)
t1 = POSTOP LEGEMBO1 %>%
 tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
  bold_p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**Leg Ischemia/Embol**")%>%as_flex_table()
## ----- adjusted -----
POSTOP_LEGEMBO2 = geeglm(POSTOP_LEGEMBO ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+
                     PRIOR_AORSURG+PRIOR_CHF+PREOP_DIALYSIS+PATHOLOGY+extent,
                   family=binomial(link="logit"), data=TEVAR_PROC,
                   id = CENTERID, corstr = "independence")
t2 = POSTOP LEGEMBO2 %>%
  tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
  bold_p(t = 0.05)
tbl_merge(tbls = list(t2),tab_spanner ="**Leg Ischemia/Embol**")%>%as_flex_table()
## ----- Post-op Respiratory -----
# table1(~ AGECAT | factor(POSTOP_RESPIRATORY), data=subset(TEVAR_PROC, !is.na(POSTOP_RESPIRATORY)))
## ----- unadjusted -----
POSTOP_RESPIRATORY1 = geeglm(POSTOP_RESPIRATORY~ PRESENTATION,
                    data=TEVAR_PROC, family=binomial(link="logit"),
                    id = CENTERID, corstr = "independence")
t1 = POSTOP_RESPIRATORY1 %>%
  tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
  bold_p(t = 0.05)
tbl_merge(tbls = list(t1),tab_spanner ="**Post-op Respiratory**")%>%as_flex_table()
## ----- adjusted -----
POSTOP_RESPIRATORY2 = geeglm(POSTOP_RESPIRATORY ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+
                     PRIOR AORSURG+PRIOR CHF+PREOP DIALYSIS+PATHOLOGY+extent,
                   family=binomial(link="logit"), data=TEVAR_PROC,
                   id = CENTERID, corstr = "independence")
t2 = POSTOP_RESPIRATORY2 %>%
  tbl_regression(exponentiate=TRUE, tidy_fun = broom.mixed::tidy,
                pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
  bold_p(t = 0.05)
tbl_merge(tbls = list(t2),tab_spanner ="**Post-op Respiratory**")%>%as_flex_table()
# ## ----- Transfusion # Units PRBC ----
```

```
# ## ----- unadjusted -----
 \# \ POSTOP\_PRBC1 < -geeglm(POSTOP\_PRBC \ \sim \ PRESENTATION, \ data = TEVAR\_PROC, id = CENTERID, \ corstr = "independence of the content of 
# t1<-POSTOP_PRBC1 %>%tbl_regression(tidy_fun = broom.mixed::tidy,
                                                                                                                        pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
#
       bold_p(t = 0.05)
#
 \# \ tbl\_merge(tbls = list(t1), tab\_spanner = "**Transfusion \# \ Units \ PRBC**") \% > \% as\_flex\_table() 
#
# ## ----- adjusted -----
# POSTOP_PRBC2<-geeglm(POSTOP_PRBC ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+PRIOR_AORSURG+PRIOR_CHF+P
# POSTOP_PRBC2<-geeglm(POSTOP_PRBC ~ PRESENTATION+AGECAT+GENDER+PREOP_SMOKING+PRIOR_AORSURG+PRIOR_CHF+P
#
#
# t2<- POSTOP_PRBC2 %>%tbl_regression(tidy_fun = broom.mixed::tidy,
#
                                                                                                                           pvalue_fun = function(x) style_pvalue(x, digits = 2))%>%
#
        bold_p(t = 0.05)
 \# \ tbl\_merge(tbls = list(t2), tab\_spanner = "**Transfusion \# \ Units \ PRBC**") \% > \% as\_flex\_table()
```