

# Confirmatory Factor Analysis

Statsomat.com

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**Warning: The automatic computation and interpretation delivered by the Statsomat should not completely replace the classical, made by humans graphical exploratory data analysis and statistical analysis. There may be data cases for which the Statsomat does not deliver the most optimal solution or output interpretation.**

## Basic Information

Automatic statistics for the file:

File
case9.csv

Your selection for the encoding: Auto

Your selection for the decimal character: Auto

Observations (rows with at least one non-missing value): 500

Variables (columns with at least one non-missing value): 7

Variables considered continuous: 5

Variables considered continuous
ACTIV
SOMA
SOCF
VITAL
GENHLTH

Numerical variables considered binary or ordinal: 2

Numerical variables considered binary or ordinal
PAIN
MENTH

**Warning: Based on an analysis of the variables considered continuous, we suspect outliers in the data. If observations are erroneous, you could drop them and restart the app. Outliers may affect negatively the execution or the results of the CFA. These are the suspected row numbers:**

Table 4: Rows With Suspected Outliers

	ACTIV	SOMA	SOCF	VITAL	GENHLTH
62	54	20	38	5	37
442	71	50	45	52	46
477	14	44	30	49	29

## Model Syntax

The following table describes the applied model equations in lavaan model syntax, either as entered by you in the text area (denoted by User=1) or established internally (User=0). The last column numbers the free parameters which are estimated.

Left hand side	Operator	Right hand side	User	Free parameter
PHYSF	=~	ACTIV	1	0
PHYSF	=~	SOMA	1	1
PHYSF	=~	PAIN	1	2
MENTF	=~	MENTH	1	0
MENTF	=~	SOCF	1	3
MENTF	=~	VITAL	1	4
GWB	=~	GENHLTH	1	0
ACTIV	~~	ACTIV	0	5
SOMA	~~	SOMA	0	6
PAIN	~~	PAIN	0	7
MENTH	~~	MENTH	0	8
SOCF	~~	SOCF	0	9
VITAL	~~	VITAL	0	10
GENHLTH	~~	GENHLTH	0	0
PHYSF	~~	PHYSF	0	11
MENTF	~~	MENTF	0	12
GWB	~~	GWB	0	13
PHYSF	~~	MENTF	0	14
PHYSF	~~	GWB	0	15
MENTF	~~	GWB	0	16

## Assumptions

Open issue

## Model Settings

## Outputs

### Model Fit Summary

lavaan 0.6-7 ended normally after 180 iterations

Estimator	ML
Optimization method	NLMINB
Number of free parameters	16

Number of observations	500
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Model Test User Model:

Test statistic	52.854
Degrees of freedom	12
P-value (Chi-square)	0.000

Model Test Baseline Model:

Test statistic	1924.279
Degrees of freedom	21
P-value	0.000

User Model versus Baseline Model:

Comparative Fit Index (CFI)	0.979
Tucker-Lewis Index (TLI)	0.962

Loglikelihood and Information Criteria:

Loglikelihood user model (H0)	-11854.592
Loglikelihood unrestricted model (H1)	-11828.165
Akaike (AIC)	23741.183
Bayesian (BIC)	23808.617
Sample-size adjusted Bayesian (BIC)	23757.832

Root Mean Square Error of Approximation:

RMSEA	0.083
90 Percent confidence interval - lower	0.061
90 Percent confidence interval - upper	0.106
P-value RMSEA <= 0.05	0.009

Standardized Root Mean Square Residual:

SRMR	0.049
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Parameter Estimates:

Standard errors	Standard
Information	Expected
Information saturated (h1) model	Structured

Latent Variables:

	Estimate	Std.Err	z-value	P(> z )
PHYSF =~				

ACTIV	1.000			
SOMA	0.857	0.046	18.773	0.000
PAIN	0.098	0.009	11.506	0.000
MENTF =~				
MENTH	1.000			
SOCF	1.853	0.072	25.770	0.000
VITAL	2.357	0.109	21.566	0.000
GWB =~				
GENHLTH	1.000			

Covariances:

	Estimate	Std.Err	z-value	P(> z )
PHYSF ~~				
MENTF	28.536	4.335	6.582	0.000
GWB	71.920	7.921	9.079	0.000
MENTF ~~				
GWB	25.160	2.282	11.024	0.000

Variances:

	Estimate	Std.Err	z-value	P(> z )
.ACTIV	111.698	14.121	7.910	0.000
.SOMA	40.660	9.300	4.372	0.000
.PAIN	7.910	0.524	15.107	0.000
.MENTH	8.582	0.752	11.408	0.000
.SOCF	11.645	1.875	6.212	0.000
.VITAL	68.304	5.270	12.961	0.000
.GENHLTH	0.000			
PHYSF	285.950	27.057	10.568	0.000
MENTF	22.107	1.935	11.425	0.000
GWB	71.462	4.520	15.811	0.000

**Completely Standardized Parameter Estimates**

Latent Variables:

	est.std	Std.Err	z-value	P(> z )	ci.lower	ci.upper
PHYSF =~						
ACTIV	0.848	0.022	38.446	0.000	0.805	0.891
SOMA	0.915	0.021	44.056	0.000	0.875	0.956
PAIN	0.509	0.036	14.112	0.000	0.438	0.580
MENTF =~						
MENTH	0.849	0.016	53.007	0.000	0.817	0.880
SOCF	0.931	0.012	76.749	0.000	0.907	0.955
VITAL	0.802	0.019	42.692	0.000	0.765	0.838
GWB =~						
GENHLTH	1.000				1.000	1.000

Covariances:

	est.std	Std.Err	z-value	P(> z )	ci.lower	ci.upper
PHYSF ~~						
MENTF	0.359	0.044	8.172	0.000	0.273	0.445
GWB	0.503	0.036	13.854	0.000	0.432	0.574
MENTF ~~						
GWB	0.633	0.029	21.863	0.000	0.576	0.690

#### Variances:

	est.std	Std.Err	z-value	P(> z )	ci.lower	ci.upper
.ACTIV	0.281	0.037	7.509	0.000	0.208	0.354
.SOMA	0.162	0.038	4.263	0.000	0.088	0.237
.PAIN	0.741	0.037	20.187	0.000	0.669	0.813
.MENTH	0.280	0.027	10.289	0.000	0.226	0.333
.SOCF	0.133	0.023	5.887	0.000	0.089	0.177
.VITAL	0.357	0.030	11.873	0.000	0.298	0.416
.GENHLTH	0.000				0.000	0.000
PHYSF	1.000				1.000	1.000
MENTF	1.000				1.000	1.000
GWB	1.000				1.000	1.000

#### Communality

Table 6: Communality

Variable	Communality
ACTIV	0.72
GENHLTH	1.00
MENTH	0.72
PAIN	0.26
SOCF	0.87
SOMA	0.84
VITAL	0.64

#### Factor Discriminant Validity

Table 7: Factor Discriminant Validity Test at Cutoff 0.85

			Factor Correlation	Chisq diff	Df diff	P-Value
PHYSF	~~	MENTF	0.359	262.289	1	<0.001
PHYSF	~~	GWB	0.503	189.921	1	<0.001
MENTF	~~	GWB	0.633	107.294	1	<0.001

#### Factor Reliability

Table 8: Factor Reliability

	PHYSF	MENTF	GWB	total
Omega (Bentler)	0.87	0.87	1	0.92
Omega (McDonald)	0.87	0.87	1	0.92
AVE	0.76	0.71	1	0.76

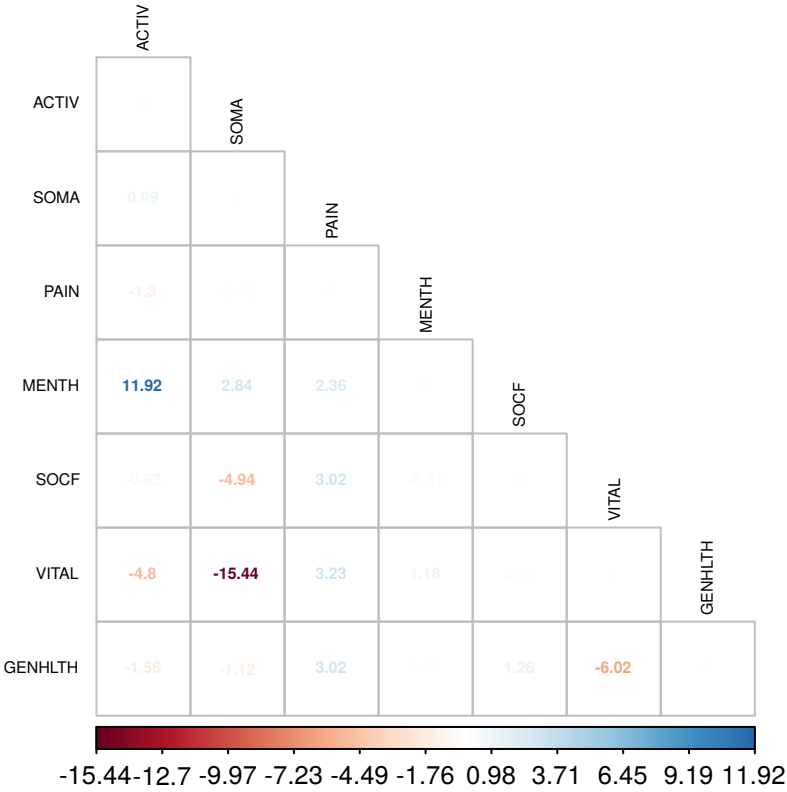
### Observed Covariance Matrix

	ACTIV						
ACTIV	397.65						
		SOMA					
SOMA	246.09	250.76					
			PAIN				
PAIN	26.82	23.95	10.68				
				MENTH			
MENTH	40.46	27.3	5.17	30.69			
					SOCF		
SOCF	52.25	40.38	8.22	40.66	87.55		
						VITAL	
VITAL	62.45	42.21	9.84	53.29	96.73	191.11	
							GENHLTH
GENHLTH	70.36	60.53	10.09	25.22	47.88	53.28	71.46

Model-Implied Covariance Matrix

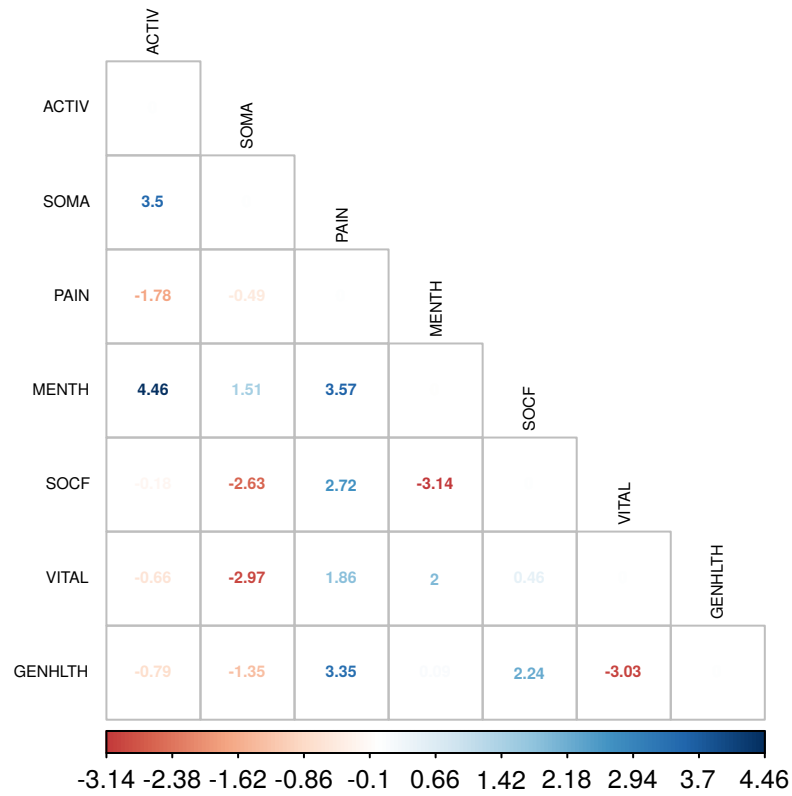
	ACTIV						
ACTIV	397.65						
		SOMA					
SOMA	245.11	250.76					
			PAIN				
PAIN	28.12	24.1	10.68				
				MENTH			
MENTH	28.54	24.46	2.81	30.69			
					SOCF		
SOCF	52.88	45.32	5.2	40.96	87.55		
						VITAL	
VITAL	67.26	57.65	6.61	52.1	96.55	191.11	
							GENHLTH
GENHLTH	71.92	61.65	7.07	25.16	46.62	59.3	71.46

Residual Covariance Matrix

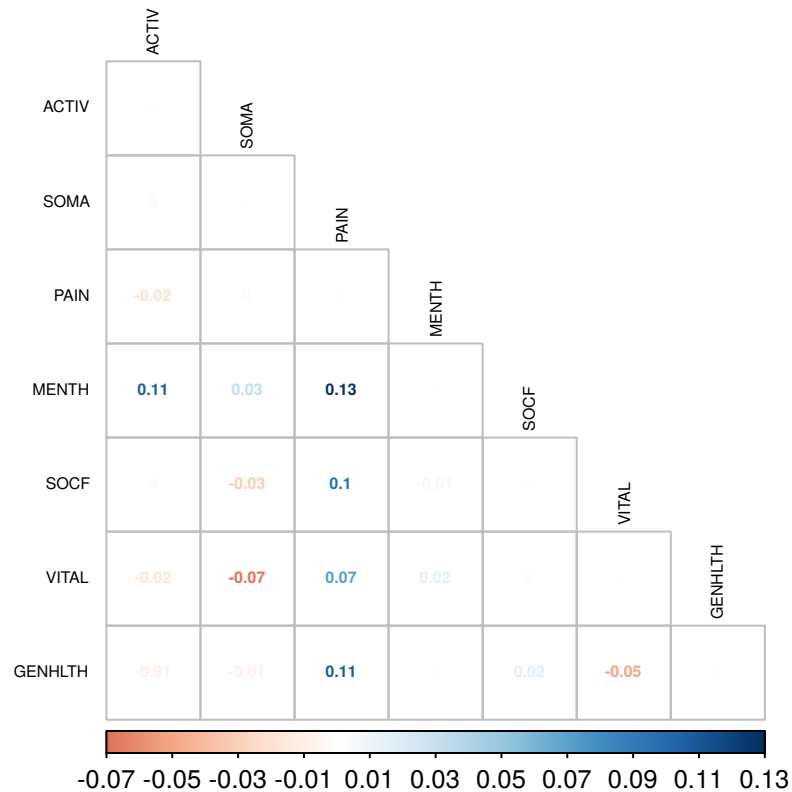




## Standardized Residual Matrix



## Residual Correlation Matrix



## Modification Indices

Table 9: Modification Indices With Respect To Error Covariances

Left	Operator	Right	Modification Index	Expected Parameter Change	Delta	Power	Decision
ACTIV	~~	MENTH	14.917	7.038	0.1	0.05	** (m) **
ACTIV	~~	SOMA	13.093	120.668	0.1	0.05	** (m) **
SOCF	~~	GENHLTH	9.961	5.367	0.1	0.05	** (m) **
MENTH	~~	SOCF	9.221	-5.967	0.1	0.05	** (m) **

Note:

Maximum 10 modification indices in descending order of their magnitude are listed.

Table 10: Modification Indices With Respect To Factor Loadings

Left	Operator	Right	Modification Index	Expected Parameter Change	Delta	Power	Decision
PHYSF	=~	MENTH	13.019	0.038	0.4	1.000	epc:nm
GWB	=~	PAIN	11.980	0.063	0.4	1.000	epc:nm
MENTF	=~	PAIN	10.697	0.102	0.4	1.000	epc:nm
GWB	=~	VITAL	8.628	-0.196	0.4	1.000	epc:nm
MENTF	=~	SOMA	8.051	-0.366	0.4	0.873	epc:nm
GWB	=~	SOCF	5.729	0.106	0.4	1.000	epc:nm

Note:

Table 10: Modification Indices With Respect To Factor Loadings (continued)

Left	Operator	Right	Modification Index	Expected Parameter Change	Delta	Power	Decision
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Maximum 10 modification indices in descending order of their magnitude are listed.

## Interpretation

### Goodness of Fit Indices

We consider some of the model fit indices from the Model Fit Summary section to check the goodness-of-fit of the model. To decide for an acceptable or non-acceptable model, we apply thresholds considered in the References: [Brown], [Kline].

#### Model Test User Model

The degrees of freedom are calculated as the number of known parameters minus the number of free parameters:  $28 - 16 = 12$ . The 12 degrees of freedom indicate an over-identified model, fact which basically enables further analysis and interpretation.

The test statistic with the value 52.854 is called the Chi-square model fit index and represents the difference between summaries of the model-implied covariance matrix and the observed covariance matrix which is hypothesized and desirable to be zero. In general, if the p-value is larger than 0.01 then the test is not statistically significant at 1 % error, the hypothesis cannot be rejected, which would be in favour of the model.

In our case, the p-value is  $< 0.001$  suggesting that the model may not be acceptable for the data. The Chi-square model fit index is based on a very stringent statistical hypothesis which may have no practical relevance. We will consider it only in connection with other model fit indices.

#### Model Test Baseline Model

The test statistic with the value 1924.279 represents the difference between summaries of the baseline model (an alternative model-implied covariance matrix having zero covariances, i.e. a worst fitting model assuming independent variables) and the observed covariance matrix. The p-value of the test of a zero difference is  $< 0.001$  suggesting that the baseline model does not fit good to the data. This result is used indirectly in the construction of other model fit indices.

#### Root Mean Square Error of Approximation:

The Root Mean Square Error of Approximation (RMSEA) is a fit index based on the chi-square test statistic, which corrects for parsimony, i.e. overly complex models are penalized. RMSEA can be greater or equal than zero, with values close to zero suggesting an acceptable model fit.

In our case, the RMSEA is 0.083. The upper bound of the 90% confidence interval of the RMSEA is 0.106 and greater or equal than the threshold value 0.1, suggesting a poor model fit.

#### Standardized Root Mean Square Residual:

The Standardized Root Mean Square Residual (SRMR) is a fit index derived from the residual correlation matrix with a range between zero and one with values close to zero suggesting an acceptable model fit.

In our case, the SRMR is 0.05 which is smaller than the threshold value 0.06 suggesting an acceptable model fit.

#### User Model versus Baseline Model

The Comparative Fit Index (CFI), evaluates the fit of the the model in relation to the worst-fitting baseline model described above. It ranges between zero and one, with values close to one suggesting good models (in the sense of

departure from the baseline model).

In our case, the CFI is 0.979 which is greater or equal than the threshold value 0.95, suggesting a good model fit.

Similarly to the CFI, the Tucker-Lewis Index (TLI) evaluates the fit of the model in relation to the worst-fitting baseline model described above. Moreover, overly complex models are penalized. Values can range outside zero and one but the index is interpreted similarly to the CFI.

In our case, the TLI is 0.962 which is greater or equal than the threshold value 0.95, suggesting a good model fit.

## Summary of the Goodness of Fit Indices

The TLI model fit index suggests an acceptable model fit. The SRMR model fit index suggests an acceptable model fit. The Chi-square model fit index and the RMSEA suggest a poor model fit. The goodness-of-fit of the model is uncertain. We proceed by diagnosing the sources of possible misspecification.

## Residuals

We analyze the residual matrices from the Outputs chapter. The residual covariance matrix represents the difference between the observed covariance matrix and the fitted model-implied covariance matrix. Large absolute values indicate local areas of misfit. However, the residuals are affected by the raw metric and are difficult to interpret more precisely.

A better interpretation allows the standardized residual matrix (residuals divided by their estimated asymptotic standard error) and the residual correlation matrix.

Following variable pairs have standardized residuals which are larger or equal than the considered threshold 2.58 [brown] or correlation residuals which are larger or equal than the considered threshold 0.1 [kline]. In these cases, the covariance relationship between the involved variables is probably underestimated:

Table 11: Pair(s) with Underestimated Covariance

Pair 1	ACTIV	MENTH
Pair 2	GENHLTH	PAIN
Pair 3	PAIN	GENHLTH
Pair 4	PAIN	MENTH
Pair 5	PAIN	SOCF
Pair 6	SOCF	PAIN

Following variable pairs have standardized residuals which are smaller or equal than the considered threshold -2.58 [brown] or correlation residuals which are smaller or equal than the considered threshold -0.1 [kline]. In these cases, the covariance relationship between the involved variables is probably overestimated:

Table 12: Pair(s) with Overestimated Covariance

Pair 1	GENHLTH	VITAL
Pair 2	MENTH	SOCF
Pair 3	SOMA	VITAL
Pair 4	VITAL	GENHLTH

Depending on the sample size, the misspecification detected by the analysis of the residual covariance resp. correlation matrices can be statistically significant but not relevant and in practice negligible. This is matter of subject in the next section(s).

## Modification Indices

In the interpretation of the modification indices table(s) we rely mostly on [brown] and [mi]. We cite from [brown]: “The modification index reflects an approximation of how much the overall model  $\chi^2$  will decrease if the fixed or constrained parameter is freely estimated.” In other words, if adding a line with a high modification index to the model, i.e. if adding a parameter, the overall goodness-of-fit may be improved. Nevertheless, this should be done only under certain conditions, described in the sequel.

We consider only modification indices greater or equal than 3.84 (which are statistically significant at 5% type I error). Next, we search only for modification indices which achieve a power of minimum 75% in detecting a (relevant) misspecification of at least 0.1 for error or factor correlations, respectively 0.4 for factor loadings. These are characterized in the decision column by the label “epc:m”. For more information with regard to the labels of the decision column, please consult the Appendix.

We remark that these conditions are not fulfilled for modification indices with respect to error covariances. Therefore, there exist no significant and relevant modification indices with respect to error covariances.

We remark that these conditions are not fulfilled for modification indices with respect to factor loadings. Therefore, there exist no significant and relevant modification indices with respect to factor loadings.

We remark that there exist no modification indices with respect to factor covariances.

## Parameter Estimates

### Factor Loadings

We remark that the completely standardized factor loadings (section “Completely Standardized Parameter Estimates”) are all statistically significant at 5% type I error. Moreover, in absolute value they are all greater than 0.4. This cutoff-value is considered in some CFA research areas a magnitude that is substantively meaningful [brown]. Please consider also cutoff-values from your particular research area when interpreting the factor loadings. We summarize the interpretation of the completely standardized factor loadings in the next table:

Table 13: Check Completely Standardized Factor Loadings

Latent Variable	Observed Variable	Loading <sup>1</sup>	P-Value	Significant? <sup>2</sup>	Relevance <sup>3</sup>	Sign <sup>4</sup>	Check
PHYSF	ACTIV	0.85	<0.001	Yes	***	—	Ok
PHYSF	SOMA	0.92	<0.001	Yes	***	—	Ok
PHYSF	PAIN	0.51	<0.001	Yes	*	—	Ok
MENTF	MENTH	0.85	<0.001	Yes	***	—	Ok
MENTF	SOCF	0.93	<0.001	Yes	***	—	Ok
MENTF	VITAL	0.80	<0.001	Yes	***	—	Ok
GWB	GENHLTH	1.00			***	—	Ok

<sup>1</sup> The completely standardized factor loading can be interpreted as the correlation with the factor.

<sup>2</sup> Completely standardized factor loading significance at 5% type I error.

<sup>3</sup> Stars correspond to factor loadings cutoff-values: 0.4, 0.6, 0.8.

<sup>4</sup> No (correct) information available. We assume the signs of the factor loadings correspond to your expectation.

<sup>5</sup> \_\_\_\_\_

Table 13: Check Completely Standardized Factor Loadings (continued)

Latent Variable	Observed Variable	Loading <sup>1</sup>	P-Value	Significant? <sup>2</sup>	Relevance <sup>3</sup>	Sign <sup>4</sup>	Check
6 _____							
7 _____							

Moreover, we remark that the significance test results for the completely standardized factor loadings from above coincide to those of the unstandardized factor loadings (within section “Model Fit Summary”, for non-marker variables).

We proceed by interpreting the (unstandardized) factor loadings from the “Model Fit Summary” section:

Table 14: Interpretation of Unstandardized Factor Loadings

Interpretation of Unstandardized Factor Loadings
A 1-unit increase in PHYSF leads to a 1.00 -unit increase in the ACTIV
A 1-unit increase in PHYSF leads to a 0.86 -unit increase in the SOMA
A 1-unit increase in PHYSF leads to a 0.10 -unit increase in the PAIN
A 1-unit increase in MENTF leads to a 1.00 -unit increase in the MENTH
A 1-unit increase in MENTF leads to a 1.85 -unit increase in the SOCF
A 1-unit increase in MENTF leads to a 2.36 -unit increase in the VITAL
A 1-unit increase in GWB leads to a 1.00 -unit increase in the GENHLTH

### Factor Discriminant Validity

As noted by [Brown], “the interpretability of the size and statistical significance of factor intercorrelations depends on the specific research context.” Though, the largest estimated factor intercorrelation within the section “Completely Standardized Parameter Estimates” is 0.63 which we regard as a proof of a reasonable discriminant validity. Moreover, the statistical test(s) for factor discriminant validity are statistically significant at 5% type I error.

### Error Variances

We summarize the interpretation of the error variances and communalities in the next table:

Table 15: Completely Standardized Error Variances and Commuality

Observed Variable	Error Variance <sup>1</sup>	Communality <sup>23</sup>	P-Value	Significant Error Variance? <sup>4</sup>
ACTIV	0.28	0.72	<0.001	Yes
GENHLTH	0.00	1.00		
MENTH	0.28	0.72	<0.001	Yes
PAIN	0.74	0.26	<0.001	Yes
SOCF	0.13	0.87	<0.001	Yes
SOMA	0.16	0.84	<0.001	Yes
VITAL	0.36	0.64	<0.001	Yes

<sup>1</sup> Can be interpreted as proportion of unexplained variance by the latent factor(s) (%).

<sup>2</sup> Corresponds to the squared factor loading.

<sup>3</sup> Can be interpreted as proportion of explained variance by the latent factor(s) (%).

<sup>4</sup> 5% type I error. Typically significant since a large portion of variance is not explained by the latent variable.

## Factor Reliability

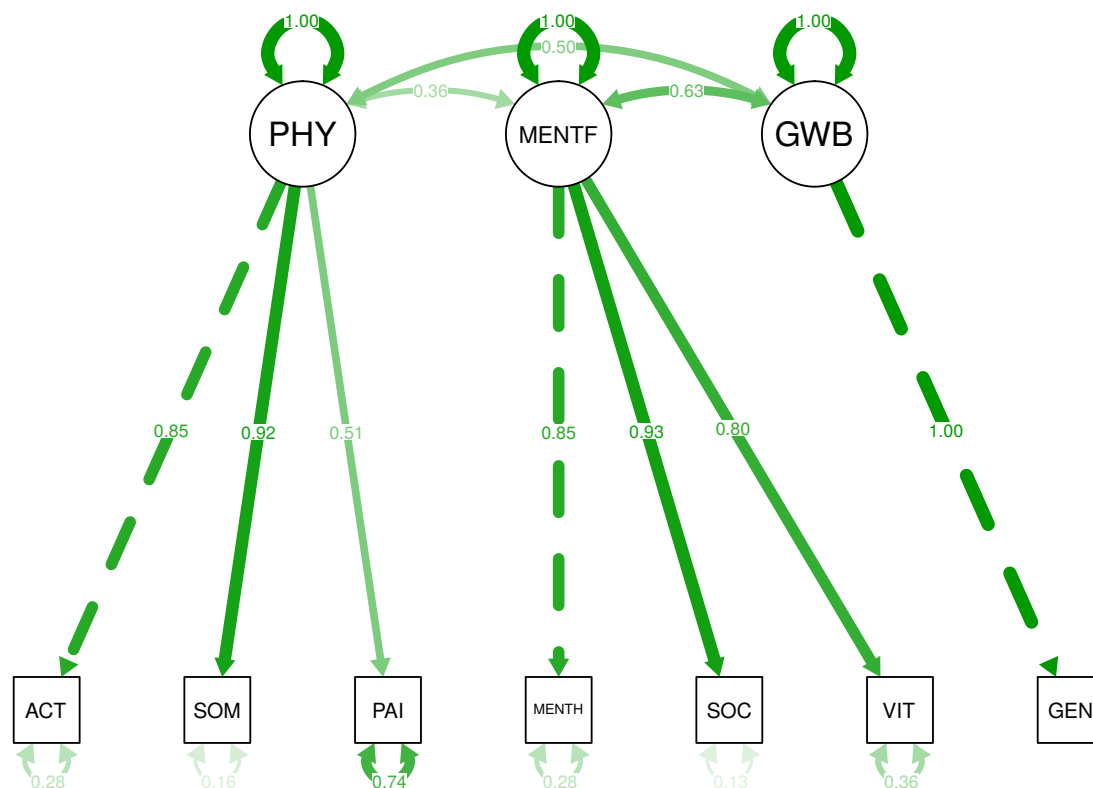
The table “Factor Reliability” contains the omega measures of factor reliability given by Bentler (Bentler, 1972, 2009) and McDonald (McDonald, 1999) and the average variance extracted (AVE). The interpretability of the reliability measures depend on the specific research context. In some fields of research, omega values greater or equal than 0.6 and AVE values greater or equal than 0.5 (fulfilled by and large in your case) could be sufficient.

## Final Summary

In our final evaluation, we distinguish between following model quality categories: acceptable, non-acceptable or uncertain.

Considering the goodness-of-fit indices, the quality of the model is uncertain. Since we cannot identify (significant and relevant) localized areas of ill fit, we assume that the model is acceptable but strongly recommend to replicate the CFA in an independent sample to validate the results. Please consider also that this summary depends on (some) hard-coded cutoff-values which may be too liberal or too conservative for your research area.

## Path Diagram



## APPENDIX

### Decision Column of the Modification Indices Table

```
not mi.significant & not high.power := "(i)"
mi.significant & not high.power := "**(m)**"
not mi.significant & high.power := "(nm)"
mi.significant & high.power & not epc.high := "epc:nm"
mi.significant & high.power & epc.high := "*epc:m*"
```

## References