

Comparison with mgcv

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```
library(tinyVAST)
library(pdp) # approx = TRUE gives effects for average of other covariates
library(lattice)
library(visreg)
set.seed(101)
```

tinyVAST is an R package for fitting vector autoregressive spatio-temporal (VAST) models using a minimal and user-friendly interface. We here show how it can replicate analysis using splines specified via `mgcv`

```
# Simulate
n_obs = 1000
x = rnorm(n_obs)
group = sample( x=1:5, size=n_obs, replace=TRUE )
w = runif(n_obs, min=0, max=2)
z = 1 + x^2 + cos((w+group/5)*2*pi) + rnorm(5)[group]
a = exp(0.1*rnorm(n_obs))
y = z + a + rnorm(n_obs, sd=0.2)
Data = data.frame( x=x, y=y, w=w, z=z, group=factor(group), a=a )

# fit model
Formula = y ~ 1 + s(group, bs="re") + poly(x, 2, raw=TRUE) + s(w, by=group, bs="ts") # + offset(a)
myfit = fit( data = Data,
             formula = Formula,
             control = tinyVASTcontrol(quiet=TRUE, trace=0) )
```

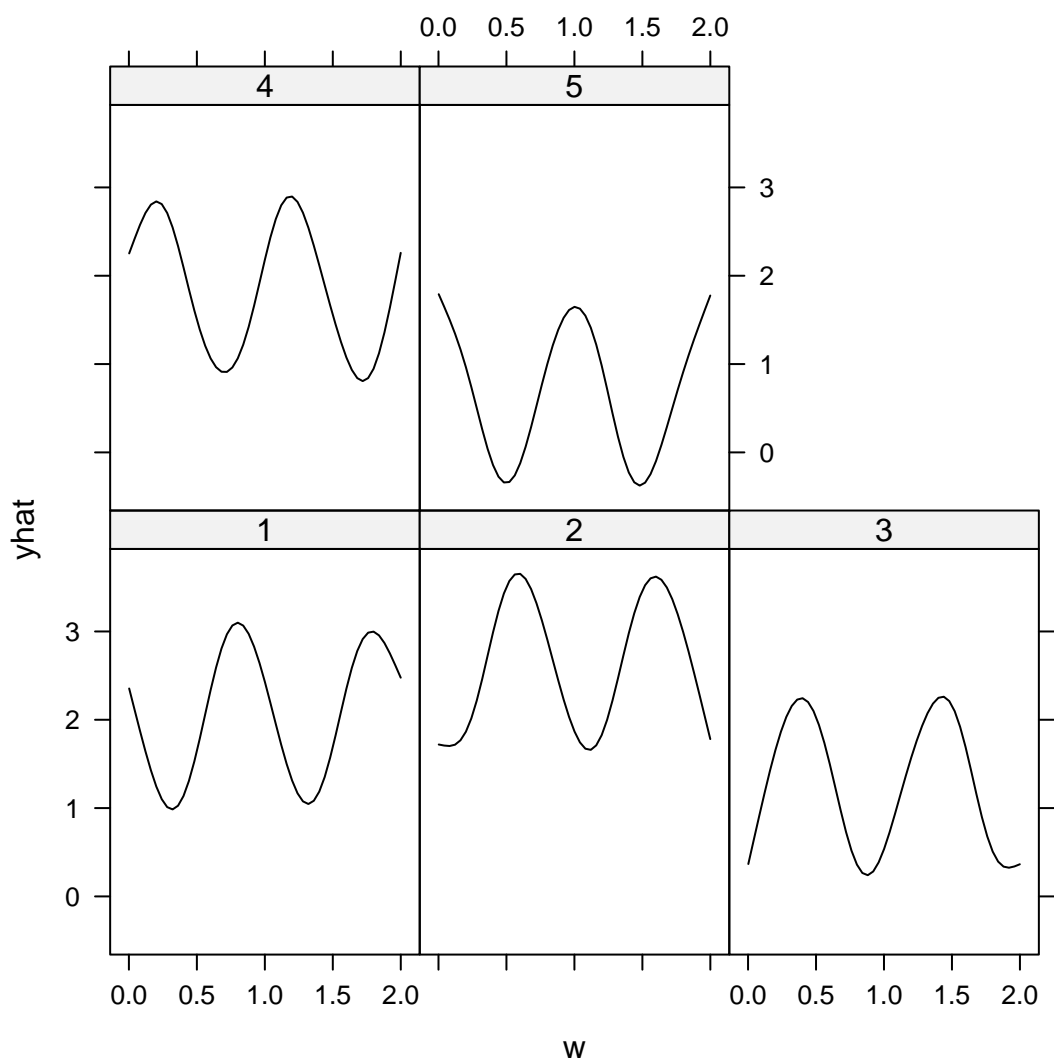
tinyVAST then has a standard predict function:

```
predict(myfit, newdata=data.frame(x=0, y=1, w=0.4, group=2, a=1) )
#> [1] 2.977754
```

and this is used to compute partial-dependence plots using package `pdp`

```
# compute partial dependence plot
Partial = partial( object = myfit,
                   pred.var = c("w", "group"),
                   pred.fun = \(object, newdata) predict(object, newdata),
                   train = Data,
                   approx = TRUE )

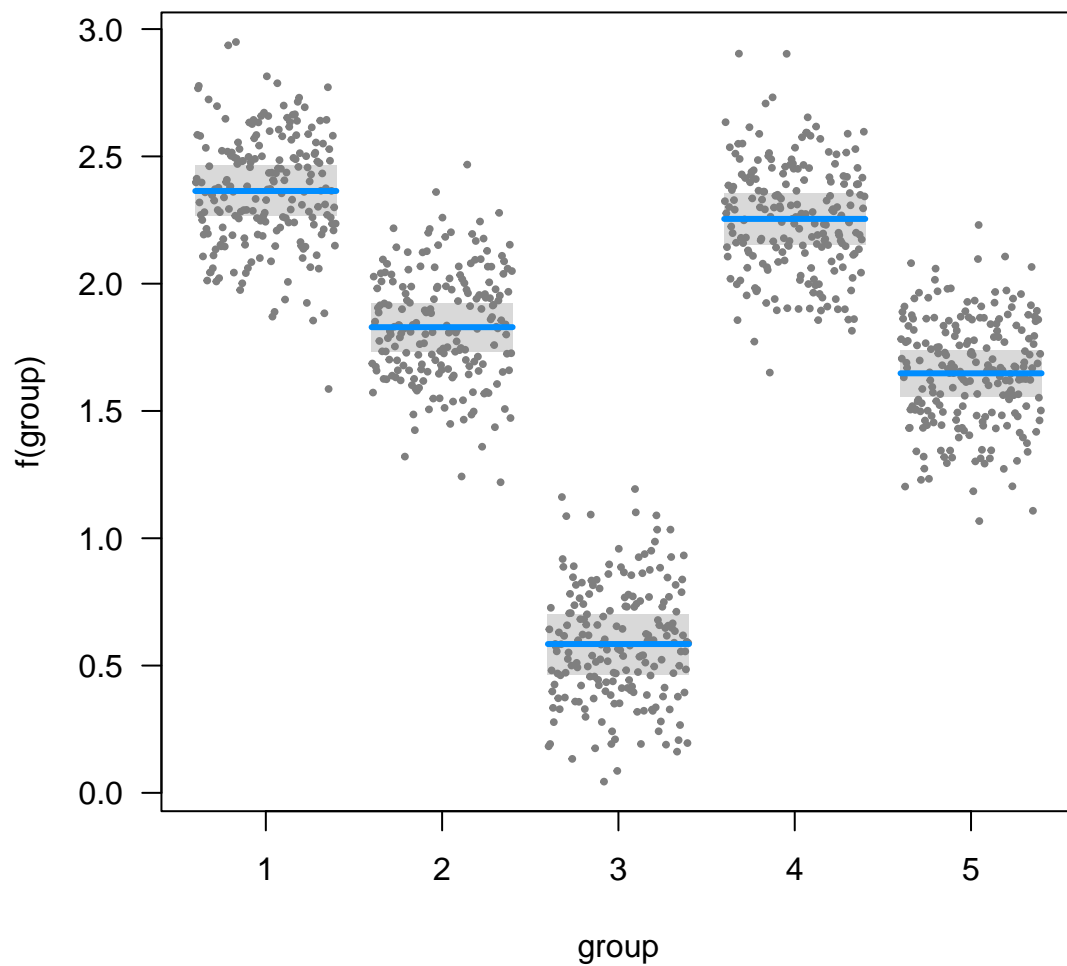
# Lattice plots as default option
plotPartial( Partial )
```



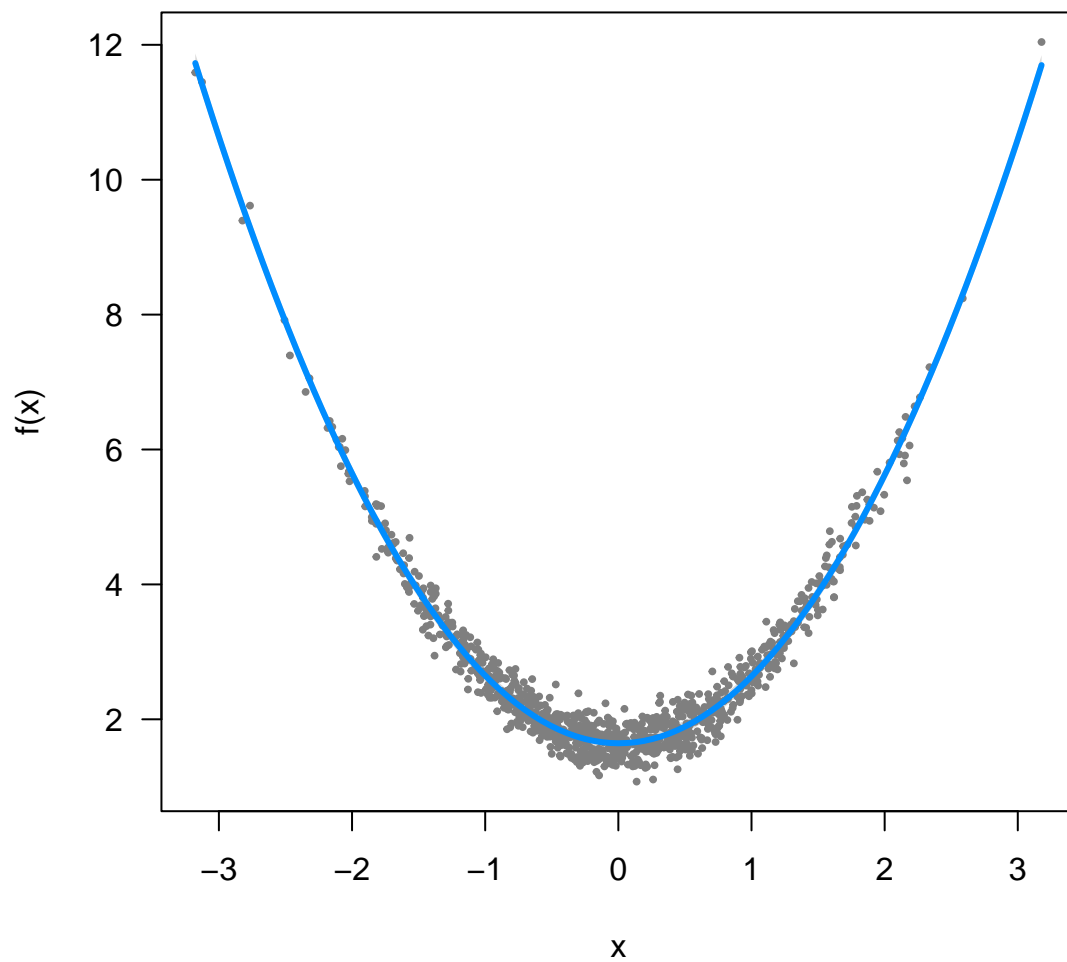
Alternatively, we can use `visreg` to visualize output:

```
visreg(myfit, xvar="group", what="p_g")
#> iter: 1  mgc: 8.184564e-13
#> Matching hessian patterns... Done
#> outer mgc: 0.001043272
#> Warning in plot.window(...): "what" is not a graphical parameter
#> Warning in plot.xy(xy, type, ...): "what" is not a graphical parameter
#> Warning in axis(side = side, at = at, labels = labels, ...): "what" is not a graphical parameter

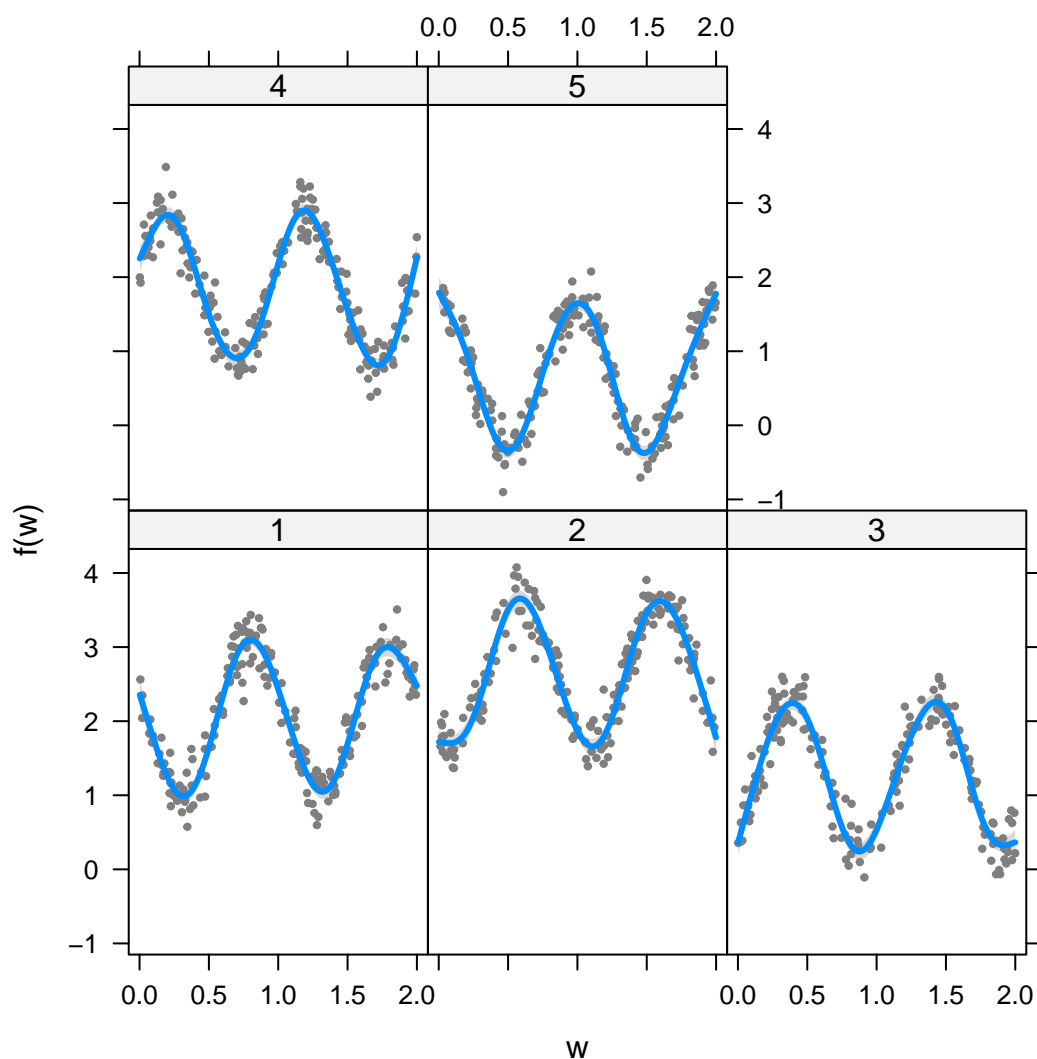
#> Warning in axis(side = side, at = at, labels = labels, ...): "what" is not a graphical parameter
#> Warning in box(...): "what" is not a graphical parameter
#> Warning in title(...): "what" is not a graphical parameter
#> Warning in axis(side = 1, at = c(0.0833333333333333, 0.291666666666667, : "what" is not a graphical parameter
```



```
visreg(myfit, xvar="x", what="p_g")
#> iter: 1   mgc: 8.184564e-13
#> Matching hessian patterns... Done
#> outer mgc: 0.001043272
#> Warning in plot.window(...): "what" is not a graphical parameter
#> Warning in plot.xy(xy, type, ...): "what" is not a graphical parameter
#> Warning in axis(side = side, at = at, labels = labels, ...): "what" is not a graphical parameter
#> Warning in axis(side = side, at = at, labels = labels, ...): "what" is not a graphical parameter
#> Warning in box(...): "what" is not a graphical parameter
#> Warning in title(...): "what" is not a graphical parameter
```



```
visreg(myfit, xvar="w", by="group", what="p_g")
#> iter: 1  mgc: 8.184564e-13
#> Matching hessian patterns... Done
#> outer mgc: 0.001043272
#> iter: 1  mgc: 8.184564e-13
#> Matching hessian patterns... Done
#> outer mgc: 0.001043272
#> iter: 1  mgc: 8.184564e-13
#> Matching hessian patterns... Done
#> outer mgc: 0.001043272
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#> Matching hessian patterns... Done
#> outer mgc: 0.001043272
#> iter: 1  mgc: 8.184564e-13
#> Matching hessian patterns... Done
#> outer mgc: 0.001043272
```



Alternatively, we can calculate derived quantities via Monte Carlo integration of the estimated density function:

```
# Predicted sample-weighted total
integrate_output(myfit)
#>           Estimate           Std. Error Est. (bias.correct) Std. (bias.correct)
#>      2616.538079           7.187831      2616.538079           NA

# True (latent) sample-weighted total
sum( Data$z )
#> [1] 1606.42
```

Similarly, we can fit a grouped 2D spline

```
# Simulate
R = exp(-0.4 * abs(outer(1:10, 1:10, FUN="-"))) )
z = mvtnorm::rmvnorm(3, sigma=kronecker(R,R) )
```

```

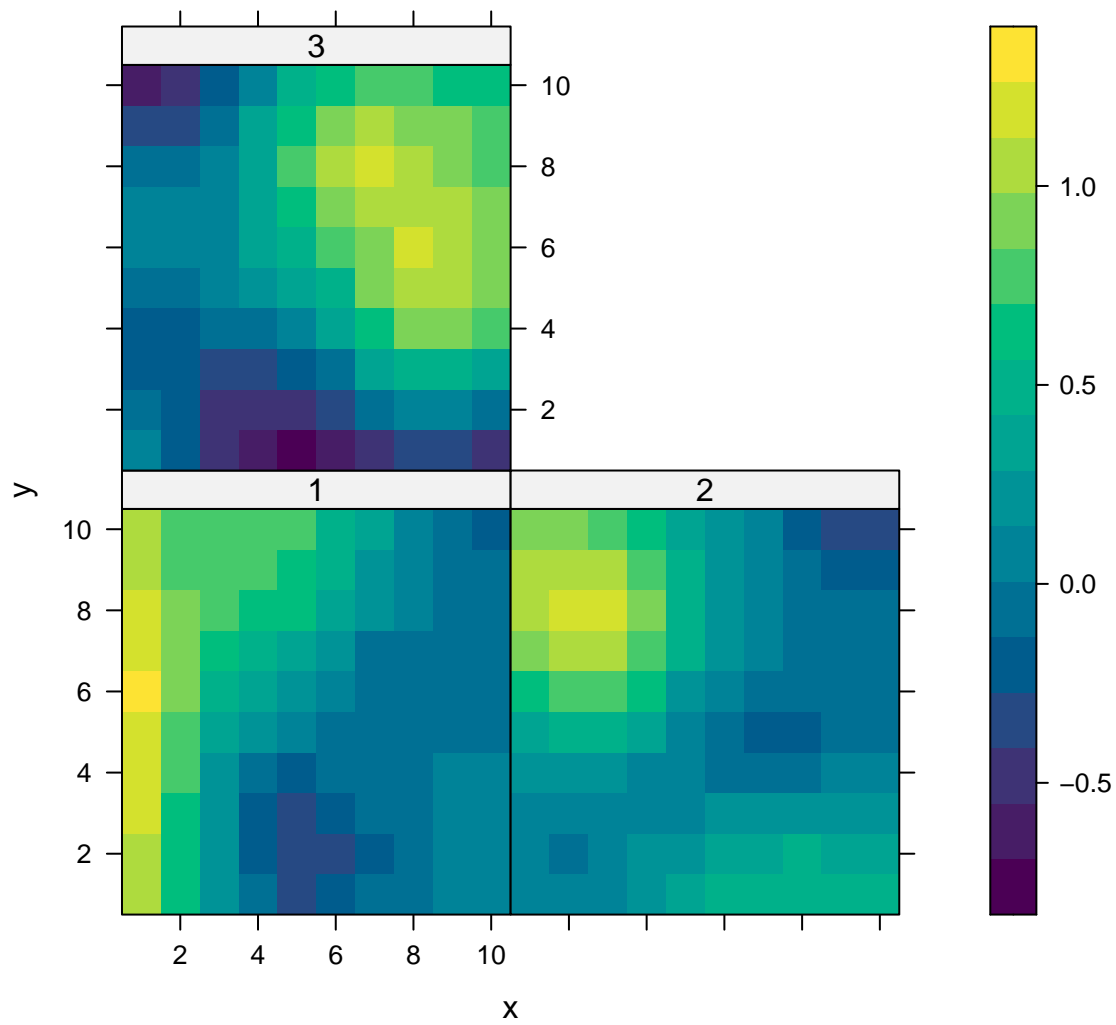
Data = data.frame( expand.grid(x=1:10, y=1:10, group=1:3), z=as.vector(t(z)))
Data$n = Data$z + rnorm(nrow(Data), sd=0.1)
Data$group = factor(Data$group)

# fit model
Formula = n ~ s(x, y, by=group)
myfit = fit( data = Data,
             formula = Formula,
             control = tinyVASTcontrol(quiet=TRUE, trace=0) )

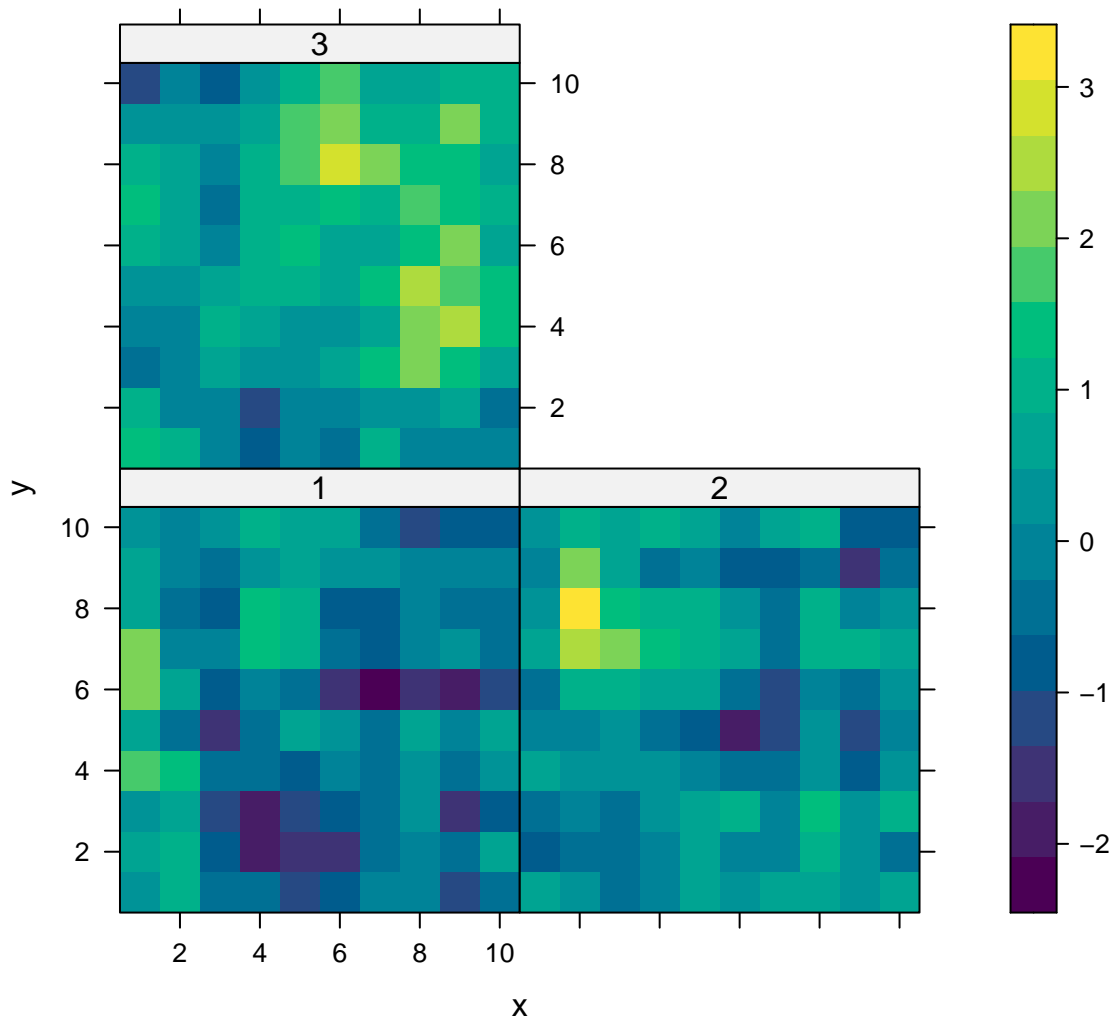
# compute partial dependence plot
mypartial = partial( object = myfit,
                     pred.var = c("x", "y", "group"),
                     pred.fun = \(object, newdata) predict(object, newdata),
                     train = Data,
                     approx = TRUE )

# Lattice plots as default option
plotPartial( mypartial )

```



```
# Lattice plot of true values
mypartial$yhat = Data$z
plotPartial( mypartial )
```



We can again use `visreg` to visualize response surfaces, although it doesn't seem possible to extract a grouped spatial term, so we here show only a single term:

```
out = visreg2d( myfit, "x", "y", cond=list("group"=1), plot=FALSE )
plot( out, main="f(x,y) for group=1")
```


$f(x,y)$ for group=1

