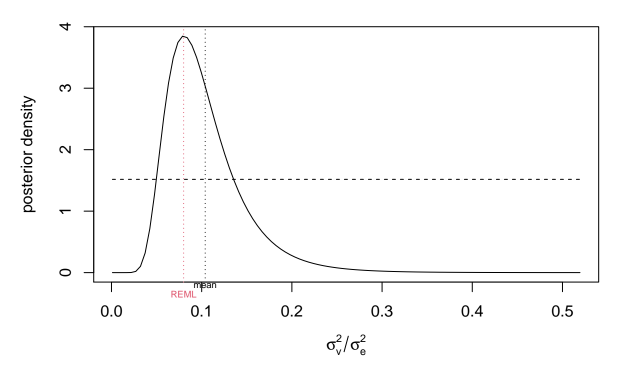
Back to hbsae

```
popdat <- m333 %>%
  group_by(subsection) %>%
  summarize(
    '(Intercept)' = n(),
    nlcd11 = mean(nlcd11))
## 'summarise()' ungrouping output (override with '.groups' argument)
popdat <- as.data.frame(popdat)</pre>
xpop <- m333 %>%
  group_by(subsection) %>%
  summarize(
   nlcd11 = mean(nlcd11)
  ) %>%
 as.data.frame()
## 'summarise()' ungrouping output (override with '.groups' argument)
rownames(xpop) <- xpop$subsection</pre>
xpop <- xpop %>% select(nlcd11)
unit_mod <- fSAE.Unit(y = m333$BIOLIVE_TPA,
                      X = m333 %>% dplyr::select(nlcd11),
                      area = m333$subsection,
                      Narea = popdat$'(Intercept)',
                      Xpop = xpop,
                      fpc = FALSE,
                      nu0 = 2,
                      s20 = 10)
## REML estimate of variance ratio: 0.08002
## numerical integration of f(x) (normalization constant): 60.69 with absolute error < 3.3e-07
## numerical integration of x*f(x): 6.305 with absolute error < 6.2e-08
## posterior mean for variance ratio: 0.1039
```

posterior density for σ_v^2/σ_e^2



```
M333Aa
                  M333Ab
                             M333Ac
##
                                         M333Ad
                                                    M333Ai
                                                               M333Ba
                                                                           M333Bb
## 0.08321000 0.06950407 0.07894010 0.09477250 0.26643264 0.03927688 0.04448655
##
       M333Bc
                  M333Ca
                             M333Cb
                                         M333Cc
                                                    M333Ce
                                                               M333Cf
## 0.13013963 0.11685367 0.04357960 0.13569412 0.11316135 0.20417707 0.17035954
                  M333Da
                             M333Db
                                         M333Dc
                                                    M333Dd
## 0.18852056 0.03178039 0.03081893 0.06035469 0.03701514 0.04089723
```

```
dat <- data.frame(
  mean = m333_area$meanbio,
  est = EST(unit_mod),
  subsection = m333_area$subsection,
  freq = EST(freq_mod)
)</pre>
```

```
dat %>%
  mutate(subsection = fct_reorder(subsection, mean)) %>%
  ggplot(aes(x = subsection)) +
  geom_point(aes(y = est,
                 color = "goldenrod"),
             position = position_nudge(x = -0.1)) +
  geom_point(aes(y = mean,
                 color = "forestgreen"),
             position = position_nudge(x = 0.1)) +
  geom_point(aes(y = freq,
                 color = "steelblue"),
             position = position_nudge(x = 0.2)) +
  theme_bw() +
  scale_color_manual(
    name = 'Estimate Type',
    values = c('goldenrod' = 'goldenrod',
               'forestgreen' = 'forestgreen',
               "steelblue" = "steelblue"),
    labels = c('Direct (Mean)', 'Hierarchical Bayesian', "BLUP"),
    guide = "legend"
  ) +
  theme(axis.text.x = element_text(
    angle = 90,
    vjust = 0.5,
   hjust = 1
  ),
  legend.position = "bottom") +
  labs(x = "Subsection",
      y = "Estimate")
```

