## 1. Page 2,

CHANGE: "This requires equating the expected proportion of genotypes shared identical by descent, given a pedigree, with the observed phenotypic correlation between relatives."

TO: "This requires equating the expected proportion of the genome shared identical by descent, given a pedigree, with the observed phenotypic correlation between relatives."

(see note0101.pdf at https://github.com/SorensenD/SLGDS)

2. Page 258, eqn. (6.1),

CHANGE second line to (brackets missing):

$$E_{y_0x} [MSE (f (x))] = E_{y_0x} [(y_0 - f (x))^2]$$

$$= E_x [E_{y_0|x} [(y_0 - f (x))^2 |x]]$$

$$= E_x [Var (y_0|x) + (E (y_0|x) - f (x))^2], \qquad (6.1)$$

3. Page 259, below eqn. (6.7),

CHANGE: "that is in the best linear approximation ..."

TO: "that is the best linear approximation ..."

4. Page 675,

CHANGE: "Boyle EA, Li YI, Pritchard JK (2017) An expanded view of complex **trits**: from polygenic to omnigenic. Cell 169:1177–1186"

TO: "Boyle EA, Li YI, Pritchard JK (2017) An expanded view of complex traits: from polygenic to omnigenic. Cell 169:1177–1186"