SMM kernel method and posterior distribution

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1 Strong duality condition

Consider a convex optimization problem

$$p^* = \min_{x \in \mathcal{D}} f_0(x) : f_i(x) \ge 0, \quad i = 1, \dots, m,$$

 $h_i(x) = 0, \quad i = 1, \dots, p,$

where the functions f_0, f_1, \ldots, f_m are convex and h_1, \ldots, h_p are affine. We can have strong duality condition through Slater's condition.

Definition 1 (Slater's condition). We say that the problem satisfies Slater's condition if it is strictly feasible, that is:

$$\exists x_0 \in \mathcal{D} : f_i(x_0) < 0, i = 1, \dots, m \quad h_i(x_0) = 0, i = 1, \dots, p$$

We can replace the above by a weak form of Slater's condition, where strict feasibility is not required whenever the function f_i is affine. We then have the

Theorem 1.1 (Slater condition Duality). If the primal problem is convex, and satisfies the weak Slater's condition, the the strong duality holds.

There are another sufficient condition for strong duality.

Theorem 1.2 (Quadratic convex optimization duality). If f_0 is quadratic convex, and the functions $f_1, \ldots, f_m, h_1, \ldots, h_p$ are all affine, then the strong duality holds, provided one of the primal or dual problems is feasible.

We can apply the second theorem to ensure the strong duality holds in our case.

2 Instability issue of the SMM algorithm

Multiple initialization can solve unstable issue of the SMM algorithm in the last meeting note. In the modified algorithm, we can set multiple initialization method. In this option, the algorithm choose the best output among multiple outputs in respect to loss function value. Figure 1 shows consistent outputs from repetitions.

3 Kernel functions for matrices

We fit the SM classifier using input feature $h(X_i)$, i = 1, ..., N. From this feature, we have the Lagrange dual problem

$$L_D = \sum_{i=1}^{N} -\frac{1}{2} \sum_{i=1}^{N} \sum_{j=1}^{N} \alpha_i \alpha_j y_i y_j \langle h(X_i), h(X_j) \rangle.$$

$$\tag{1}$$

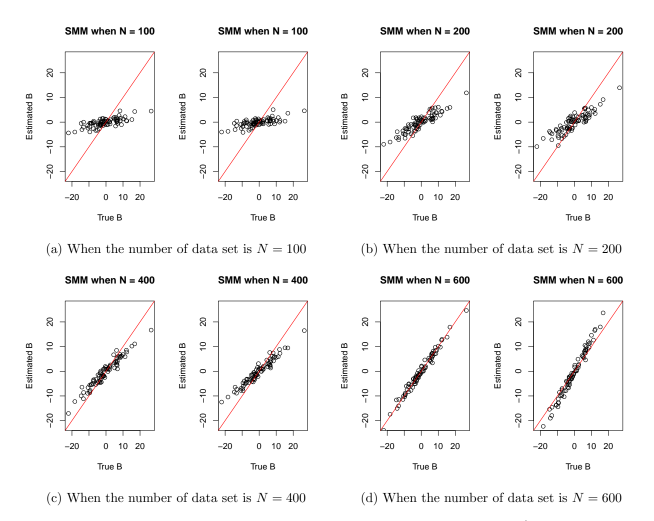


Figure 1: True parameter B is compared with multiple initialized SMM result \hat{B} under the several number of data sets $N \in \{100, 200, 400, 600\}$. The horizontal axis is entries of \hat{B} and the vertical axis is entries of \hat{B} . The number of initialization is 10. For each sub figure, we can check that the outputs are pretty much the same.

By solving (1), we obtain the nonlinear function $\hat{f}(X) = \sum_{i=1}^{N} \alpha_i y_i \langle h(X), h(X_i) \rangle$. Since all related equations require only knowledge of the kernel function,

$$K(X, X') = \langle h(X), h(X') \rangle.$$

Our goal is to define the kernel function which catches matrix structure well. In SVM case, three popular choices for K are

dth-Degree polynomial :
$$K(\boldsymbol{x}, \boldsymbol{x}') = (1 + \langle \boldsymbol{x}, \boldsymbol{x}' \rangle)^d$$
,
Radial basis : $K(\boldsymbol{x}, \boldsymbol{x}') = \exp(-\gamma \|\boldsymbol{x} - \boldsymbol{x}'\|^2)$,
Sigmoid : $K(\boldsymbol{x}, \boldsymbol{x}') = \tanh(\gamma_1 \langle \boldsymbol{x}, \boldsymbol{x}' \rangle + \gamma_2)$.

I define two measures to generalize $\langle x, x' \rangle$ and $||x - x'||^2$ into matrices case not vectorizing matrices. For two matrices $X, X' \in \mathbb{R}^{m \times n}$, we have singular value decomposition of two matrices.

$$X = \sum_{k=1}^{m \vee n} \sigma_k u_k v_k^T \quad \text{and} \quad X' = \sum_{k=1}^{m \vee n} \sigma_k' u_k' (v_k')^T.$$

From this notation, I define weighted inner product between two matrices.

$$\langle X, X' \rangle_M = \sum_{k=1}^{m \vee n} \sigma_k \sigma'_k \langle u_k, u'_k \rangle \langle v_k, v'_k \rangle. \tag{2}$$

In (2), $\sigma\sigma'$ works as weight on principal inner products of subspace and $\langle u, u' \rangle, \langle v, v' \rangle$ represent principal inner product in column space and row space respectively. From this new definition, we can generalize d-th degree polynomial kernel and sigmoid kernel into matrices case.

dth-Degree polynomial :
$$K(X, X') = (1 + \langle X, X' \rangle_M)^d$$
,
Sigmoid: $K(X, X') = \tanh(\gamma_1 \langle X, X' \rangle_M + \gamma_2)$.

In addition to inner product, we can define weighted matrices distance as

$$||X - X'||_M^2 = \sum_{k=1}^{m \vee n} \sigma_k \sigma_k' (||u_k - u_k'||^2 + ||v_k - v_k'||^2).$$
(3)

In (3), $\sigma\sigma'$ works as weight on principal row and column distances. $||u_k - u_k'||^2$ and $||v_k - v_k'||^2$ are column-wise and row-wise distances between principal vectors. With this definition we define generalized Radial basis kernel as

Radial basis :
$$K(X, X') = \exp(-\gamma ||X - X'||^2)$$
.

If two vectors x, x' are expressed as

$$x = \frac{x}{\|x\|} \|x\| \cdot 1$$
 and $x' = \frac{x'}{\|x'\|} \|x'\| \cdot 1$

We can check those definitions are consistent to vector case as follows

$$egin{aligned} \langle oldsymbol{x}, oldsymbol{x}'
angle_M &= \|oldsymbol{x} \| \|oldsymbol{x}' \| \langle oldsymbol{x}' \| \| oldsymbol{x}' \|_M &= \|oldsymbol{x} \| \| oldsymbol{x}' \| \left(\left\| \frac{oldsymbol{x}}{\| oldsymbol{x} \|} - \frac{oldsymbol{x}'}{\| oldsymbol{x}' \|}
ight\|_2^2 + \|1 - 1\|
ight) = \|oldsymbol{x} - oldsymbol{x}' \| oldsymbol{x} \| oldsymbol{x} - oldsymbol{x}' \| oldsymbol{x} - oldsymbol{x} - oldsymbol{x}' \| oldsymbol{x} - oldsymbol$$

4 Weighted binary classification

To obtain posterior distribution given feature data, we solve the regularization problem based on the weighted hinge loss.

$$\min_{\boldsymbol{\beta},\boldsymbol{\xi}} \frac{1}{2} \|\boldsymbol{\beta}\|^2 + C \left[(1-\pi) \sum_{y_i=1} \xi_i + \pi \sum_{y_i=-1} \xi_i \right]
\text{subject to } y_i(\langle x_i, \boldsymbol{\beta} \rangle + b) > 1 - \xi_i,$$
(4)

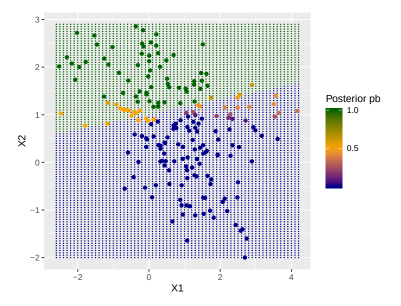


Figure 2: The green area is labeled as 1 with the SVM and blue area is -1. We can obtain non trivial posterior probability around the classification boundary.

$$\xi_i \geq 0$$
.

The related dual problem for (4) is

$$\max_{\alpha} \sum_{i=1}^{N} \alpha_{i} - \frac{1}{2} \sum_{i=1}^{N} \sum_{j=1}^{N} \alpha_{i} \alpha_{j} y_{i} y_{j} \langle \boldsymbol{x}_{i}, \boldsymbol{x}_{j} \rangle,$$
subject to $0 \leq \alpha_{i} \leq C(1-\pi)$ for $y_{i} = 1$,
$$0 \leq \alpha_{i} \leq C\pi \text{ for } y_{i} = -1,$$

$$\sum_{i=1}^{N} \alpha_{i} y_{i} = 0.$$
(5)

From the solution of (5), we can find the primal solution as $\beta = \sum_{i=1}^{N} y_i \alpha_i x_i$. With this relation, smm in R-codes section solves the equation (4). Figure 3 shows the weighted hinge loss SVM classifier with $\pi \in \{0.001, 0.5, 0.999\}$. It is known that the minimizer $\operatorname{sign}(f_{\pi}(x))$ to Equation (4) is a consistent estimate of $\operatorname{sign}(\mathbb{P}(y=1|x)-\pi)$. Therefore solving Equation (4) using different π values such that $\pi_1 < \cdots < \pi_m$, we can estimate

$$\hat{\mathbb{P}}(y=1|x) = \frac{1}{2} \left(\arg\max_{\pi_j} \{ sign(f_{\pi_j}(x)) = 1 \} + \arg\max_{\pi_j} \{ sign(f_{\pi_j}(x)) = -1 \} \right).$$
 (6)

Figure 2 shows the posterior probability estimation with the rule of (6).

5 One issue for posterior estimation

I found one issue to estimate posterior probability $\mathbb{P}(y=1|\mathbf{x})$. There are some points \mathbf{x}_i 's such that $\operatorname{sign}(\mathbb{P}(y|\mathbf{x}_i)-\pi)$ is not decreasing in respect to π . We can check that the red point in Figure 3

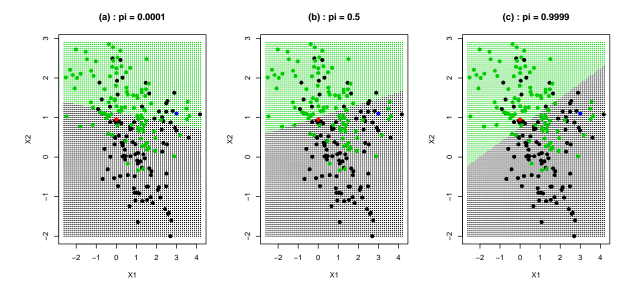


Figure 3: The sub figures show weighted hinge loss SVM classifier when $\pi = 0.001, 0.5, 0.999$. Green points represent for y = 1 and black for y = -1. The middle sub figure ($\pi = 0.5$) is the regular SVM. The red point changes its label from -1 to 1 as π increases. The blue point shows vice versa.

has $\operatorname{sign}(\mathbb{P}(y=1|\boldsymbol{x})-0.0001)=-1$ but $\operatorname{sign}(\mathbb{P}(y=1|\boldsymbol{x})-0.9999)=1$ which does not make sense. This phenomenon happens in all points located below classification boundary when $\pi=0.0001$ and above the boundary when $\pi=0.9999$ at the same time. In addition, this area is inevitable unless two classification boundaries are parallel which is hard to be satisfied. If I stick to the rule in (6), all points in the area has 0.5 as posterior probability.

6 R-codes

6.1 Updated functions

```
1 library(pracma)
  library(quadprog)
  eps = 10^{-5}
  objv = function(B, b0, X, y, cost = 10, prob = F){
    if (prob == F) {
      value = sum(B*B)/2+cost*sum(pmax(1-y*unlist(lapply(X,function(x) sum(B*x)+b0))
      ,0))
    }else{
      ind = which(y==1)
      value = sum(B*B)/2 +
13
         (1-prob)*cost*sum(pmax(1-y[ind]*unlist(lapply(X[ind],function(x) sum(B*x)+b0
14
        prob*cost*sum(pmax(1-y[-ind]*unlist(lapply(X[-ind],function(x) sum(B*x)+b0))
      ,0))
16
17
    return (value)
```

```
19 }
20
21 # Generating dataset
22 gendat = function(m,n,r,N,b0){
    result = list()
    # simulation
24
    # Weight
25
    rU = matrix(runif(m*r,-1,1),nrow = m)
    rV = matrix(runif(n*r,-1,1),nrow = n)
27
    B = rU%*%t(rV)
    # predictor matrix
30
    X = list()
31
    for (i in 1:N) {
32
     X[[i]] \leftarrow matrix(runif(m*n,-1,1),nrow = m,ncol=n)
33
34
35
    # classification
36
    v = list()
37
    for (i in 1:N) {
38
      y[[i]] = sign(sum(B*X[[i]])+b0)
39
40
41
    y = unlist(y)
42
    # predictor vector
43
    x = matrix(nrow = N, ncol = m*n)
44
    for(i in 1:N){
45
     x[i,] = as.vector(X[[i]])
46
47
    dat = data.frame(y = factor(y), x)
48
49
    result$B = B
50
    result$X = X; result$y = y; result$dat = dat
51
    return(result)
52
53 }
54
sernelm = function(X,H,y,type = c("u","v")){
    n = length(X)
57
    x = matrix(unlist(X), nrow = length(X), byrow = T)
58
    if (type == "u") {
59
     hx = matrix(unlist(lapply(X, function(x) x%*%H)), nrow = length(X), byrow = T)
60
61
      hx = matrix(unlist(lapply(X,function(x) H%*%x)),nrow = length(X),byrow = T)
62
63
    Q = matrix(nrow = n,ncol = n)
64
    for (i in 1:n) {
65
      for(j in i:n){
66
         Q[i,j] = sum(x[i,]*hx[j,])*y[i]*y[j]
67
68
         Q[j,i] = Q[i,j]
      }
69
    }
70
    h = eigen(Q)
71
    Q = (h$vectors)%*%diag(pmax(h$values,eps))%*%t(h$vectors)
    return(Q)
74 }
75
76
77
```

```
78
79 ## SMM with multiple initialization
   smm = function(X,y,r,cost = 10,rep = 10){
81
     result = list()
82
     # SMM
83
     m = nrow(X[[1]]); n = ncol(X[[1]]); N = length(X)
84
85
     compareobj = 10^100
86
     for (i in 1:rep) {
88
       error = 10
89
       iter = 0
       #initialization
90
       U = randortho(m)[,1:r]
91
       # U = matrix(runif(m*r,-1,1),nrow = m)
92
93
       V = randortho(n)[,1:r]
       # V = matrix(runif(n*r,-1,1),nrow = n)
94
       obj = objv(U%*%t(V),0,X,y,cost);obj
95
96
       while ((iter <20) & (error >10^-3)) {
97
         # update U fixing V
98
         Vs = V%*%solve(t(V)%*%V)
         H = Vs\%*\%t(V)
         dvec = rep(1,length(X))
         Dmat = kernelm(X,H,y,"u")
         Amat = cbind(y, diag(1, N), -diag(1, N))
103
         bvec = c(rep(0,1+N), rep(-cost,N))
104
         alpha = solve.QP(Dmat, dvec, Amat, bvec, meq =1)
105
         Bpart=matrix(t(y*alpha$solution)%*%matrix(unlist(X),nrow = length(X),byrow =
106
        T), nrow = m)
         U = Bpart%*%Vs
107
108
109
         # update V fixing U
110
         Us = U%*%solve(t(U)%*%U)
         H = Us\%*\%t(U)
113
         Dmat = kernelm(X,H,y,"v")
         alpha = solve.QP(Dmat, dvec, Amat, bvec, meq = 1)
114
         Bpart=matrix(t(y*alpha$solution)%*%matrix(unlist(X),nrow = length(X),byrow =
115
        T), nrow = m)
         V = t(Bpart)%*%Us
116
117
118
         ## intercept estimation
119
         Bhat = U%*%t(V); Bhat
120
         positiv = min(unlist(lapply(X,function(x) sum(Bhat*x)))[which(y==1)])
         negativ = max(unlist(lapply(X,function(x) sum(Bhat*x)))[which(y==-1)])
         if ((1-positiv)<(-1-negativ)) {</pre>
123
            b0hat = -(positiv+negativ)/2
         }else{
            gridb0 = seq(from = -1-negativ, to = 1-positiv, length = 100)
126
            b0hat = gridb0[which.min(sapply(gridb0,function(b) objv(Bhat,b,X,y)))]
128
         obj = c(obj,objv(Bhat,b0hat,X,y,cost));obj
130
         iter = iter + 1
         error = abs(-obj[iter+1]+obj[iter])/obj[iter];error
132
       if (compareobj>obj[iter+1]) {
```

```
compareobj = obj[iter+1]
136
         predictor = function(x) sign(sum(Bhat*x)+b0hat)
137
         result$B = Bhat; result$b0 = b0hat; result$obj = obj; result$iter = iter
138
         result$error = error; result$predict = predictor
139
140
     }
141
142
     return(result)
143 }
144
145
146 kernelmat = function(x,y,kernels = function(x1,x2) sum(x1*x2)){
147
     N = length(y)
     Q = matrix(nrow = N,ncol = N)
148
     for (i in 1:N) {
149
150
       for(j in i:N){
         Q[i,j] =kernels(x[i,],x[j,])*y[i]*y[j]
151
         Q[j,i] = Q[i,j]
152
153
     }
154
155
     h = eigen(Q)
     Q = (h$vectors)%*%diag(pmax(h$values,eps))%*%t(h$vectors)
156
     return(Q)
158 }
159
160
161 # SVM with kernel functions and weighted cost function
162 svm = function(X,y,cost = 10, kernels = function(x1,x2) sum(x1*x2), p = .5){
     if (p==.5) {
       cost = 2*cost
164
165
     result = list()
166
     error = 10
167
     iter = 0
168
169
     # SVM
     m = nrow(X[[1]]); n = ncol(X[[1]]); N = length(X)
170
     x = matrix(unlist(X), nrow = N, byrow = T)
     dvec = rep(1,length(X))
173
     Dmat = kernelmat(x,y,kernels)
174
     Amat = cbind(y, diag(1, N), -diag(1, N))
175
     bvec = c(rep(0,1+N), ifelse(y==1,-cost*(1-p),-cost*p))
176
     alpha = solve. QP (Dmat, dvec, Amat, bvec, meq =1)
177
178
     Bhat=matrix(t(y*alpha$solution)%*%x,nrow = m)
179
     b0hat = -(min(unlist(lapply(X,function(x) sum(Bhat*x)))[which(y==1)])+
180
                  max(unlist(lapply(X,function(x) sum(Bhat*x)))[which(y==-1)]))/2
181
     obj = objv(Bhat,b0hat,X,y,cost,prob = p)
182
184
     predictor = function(x) sign(sum(Bhat*x)+b0hat)
     result$B = Bhat; result$b0 = b0hat; result$obj = obj;
185
     result$predict = predictor
186
     return(result)
187
188
189
190
191
192
posterior = function(X,y,cost = 10,test,kernels = function(x1,x2) sum(x1*x2)){
```

```
a = 1:99
194
195
     for(i in 1:99){
        fit = svm(X,y,cost, kernels, p = i*0.01) $predict
196
197
        a[i] = fit(test)
198
     if (all(a==1)) {
199
       return(1)
200
     }else if(all(a==-1)){
201
       return(0)
202
     }else{
        return ((\max(\text{which}(a==1))+\min(\text{which}(a==-1)))/200)
204
205
206 }
```

6.2 Simulations

```
load(file = "ESL.mixture.rda")
2 names (ESL.mixture)
3 \text{ rm}(x,y)
4 attach(ESL.mixture)
5 y = ifelse(y==1,1,-1)
6 X = lapply(seq_len(nrow(x)), function(i) x[i,,drop = F])
7 \text{ par}(\text{mfrow} = c(1,1))
8 plot(x, col = y + 3)
9 dat = data.frame(y = factor(y), x)
11
12
13 a1 = matrix(nrow = 2, ncol = 99)
a1[1,] = (1:99)*0.01
15 for (i in 1:99) {
fit = svm(X,y,cost, kernels, p = i*0.01) predict
    a1[2,i] = fit(test)
17
18 }
19
20 a2 = matrix(nrow = 2, ncol = 99)
a2[1,] = (1:99)*0.01
22 for(i in 1:99){
  fit = svm(X,y,cost, kernels, p = i*0.01) $predict
23
    a2[2,i] = fit(test2)
24
25 }
26
27 a1
29
31 ### Changing svm according to weight
_{32} par(mfrow = _{c}(1,3))
34 fit = svm(X,y,cost = 10,p=0.0001) $predict
ss xgrid = expand.grid(X1 = px1, X2 = px2)
36 ygrid = apply(xgrid,1,fit)
37 plot(xgrid, col = as.numeric(ygrid+2), pch = 20, cex = .2,main = "(a) : pi =
      0.0001")
_{38} points(x, col = y + 2, pch = 19)
39 points(test, col = 'red', pch = 19)
40 points(test2, col = 'blue', pch = 19)
42 fit = svm(X,y,cost = 10,p=0.5) $predict
```

```
43 xgrid = expand.grid(X1 = px1, X2 = px2)
44 ygrid = apply(xgrid,1,fit)
45 plot(xgrid, col = as.numeric(ygrid+2), pch = 20, cex = .2,main = "(b) : pi = 0.5")
46 points(x, col = y + 2, pch = 19)
47 points(test, col = 'red', pch = 19)
48 points(test2, col = 'blue', pch = 19)
49
50
fit = svm(X,y,cost = 10,p=0.9999)$predict
xgrid = expand.grid(X1 = px1, X2 = px2)
53 ygrid = apply(xgrid,1,fit)
54 plot(xgrid, col = as.numeric(ygrid+2), pch = 20, cex = .2,main = "(c) : pi =
      0.9999")
points(x, col = y + 2, pch = 19);
56 points(test, col = "red", pch = 19)
57 points(test2,col = "blue",pch = 19)
59
60
61 ### posterior
62 posterior(X,y,cost = 10,test)
63 posterior(X,y,cost = 10,test2)
64 yposterior = vector(length = 200)
65 for(i in 1:200){
66
    yposterior[i] = posterior(X,y,cost = 10,x[i,])
   print(paste(i, "th point is done lol"))
67
68 }
69 ypost = ifelse(yposterior == -1, 0, yposterior)
70
72 \text{ par}(\text{mfrow} = c(1,1))
73 xgrid = expand.grid(X1 = px1, X2 = px2)
74 ygrid = apply(xgrid,1,fit)
75 datgrid = cbind(ygrid, xgrid)
77 \text{ colnames}(x) = c("X1", "X2")
78 realdat =as.data.frame(cbind(ypost,x))
79 breaks <-c(0.5,1,3.2)
ggplot(data = datgrid,aes(x = X1,y= X2,colour = ifelse(ygrid==-1,0,ygrid)))+geom_
      point(size = 0.001) +
    geom_point(data = realdat,aes(x = realdat[,2],y = realdat[,3],colour = ypost))+
81
    labs(colour = "Posterior pb")+
    scale_colour_gradientn(colours = c("darkblue","orange","darkgreen"),
                                                           breaks = breaks, labels =
84
   format(breaks))
```