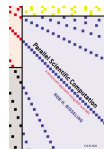
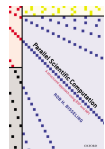


Program `bspfft` (PSC §3.6)



Sequential unordered FFT: specification

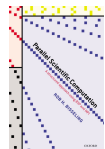
```
void ufft(double *x, int n, int sign, double *w) {  
  
    /* This sequential function computes the  
       unordered DFT of a complex vector x of  
       length n,  $n=2^m$ ,  $m \geq 0$ .  
       If sign = 1, the forward DFT  $FRx$  is computed.  
       If sign = -1, the backward UDFT  $\text{conjg}(F)Rx$ ,  
       where F is the n by n Fourier matrix and  
           R the n by n bit-reversal matrix.  
       The output overwrites x.  
       w is a table of n/2 complex weights,  
            $\exp(-2\pi i \cdot j/n)$ ,  $0 \leq j < n/2$ ,  
       which must have been initialised before  
       calling this function.  
    */
```



Data structure

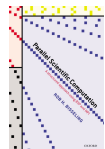
- A complex vector \mathbf{x} of length n is stored as a real array \mathbf{x} of size $2n$.
- The real and imaginary parts alternate:

$$\begin{aligned} \mathbf{x}[2 * j] &= \text{Re}(x_j) \\ \mathbf{x}[2 * j + 1] &= \text{Im}(x_j) \end{aligned}$$



Sequential unordered FFT: body

```
for(k=2; k<=n; k *=2){
    nk= n/k;
    for(r=0; r<nk; r++){
        rk= 2*r*k;
        for(j=0; j<k; j +=2){
            wr= w[j*nk]; // exp(-2*pi*i*j*(n/k)/n)
                        // = exp(-2*pi*i*j/k)
            if (sign==1)
                wi= w[j*nk+1];
            else
                wi= -w[j*nk+1];
            j0= rk+j; j1= j0+1;
            j2= j0+k; j3= j2+1;
            taur= wr*x[j2] - wi*x[j3];
            taui= wi*x[j2] + wr*x[j3];
            x[j2]= x[j0]-taur;    x[j3]= x[j1]-taui;
            x[j0] += taur;        x[j1] += taui;
        }
    }
}
```



Permutation to be used for bit reversal $\sigma = \rho_n$

```
void permute(double *x, int n, int *sigma){  
    /* This sequential function permutes a  
       complex vector x by the permutation sigma  
       (decomposable into disjoint swaps),  
       y[j] = x[sigma[j]], 0 <= j < n.  
       The output overwrites the vector x. */
```

```
    int j, j0, j1, j2, j3;  
    double tmpr, tmpi;
```

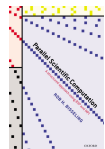
```
    for(j=0; j<n; j++){  
        if (j<sigma[j]){ // swap j and sigma[j]  
            j0= 2*j;           j1= j0+1;  
            j2= 2*sigma[j];    j3= j2+1;  
            tmpr= x[j0];       tmpi= x[j1];  
            x[j0]= x[j2];      x[j1]= x[j3];  
            x[j2]= tmpr;       x[j3]= tmpi;
```



Initialisation of bit reversal $\rho_n, n = 2^m \geq 2$

```
void bitrev_init(int n, int *rho){
    int j;
    unsigned int n1=n, rem, val, k, lastbit, one=1;

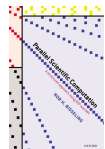
    for(j=0; j<n; j++){
        rem= j; // j= (b(m-1),...,b1,b0) in binary
        val= 0;
        for (k=1; k<n1; k <= 1){
            lastbit= rem & one;
            // lastbit = b(i) with i= log2(k)
            rem >>= 1; // rem = (b(m-1),...,b(i+1))
            val <<= 1;
            val |= lastbit; // val = (b0,...,b(i))
        }
        rho[j]= (int)val;
    }
}
```



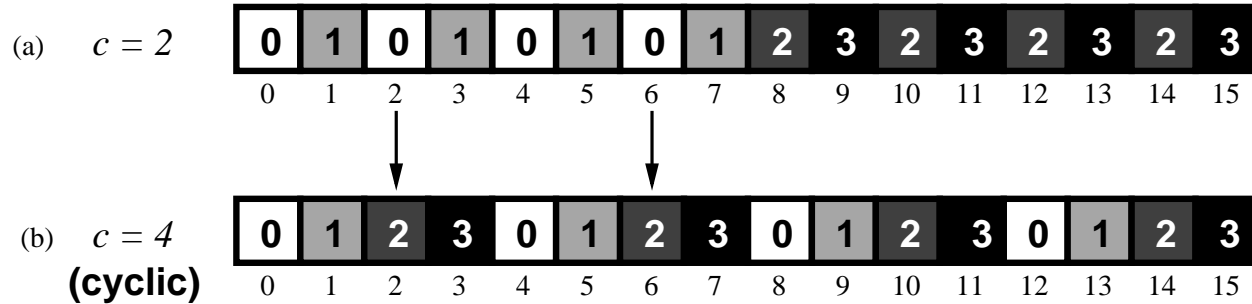
Assertions inside a loop (**loop invariants**) are powerful!

Bit operations

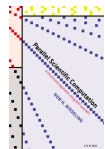
- Rule: use bit operations only sparingly in scientific computation.
- Reason: they obfuscate code, and good compilers often make them unnecessary.
- Here we encounter an exception: the cost of the bit reversal is of the same order $\mathcal{O}(n \log_2 n)$ as that of the FFT itself, so the bit reversal is important, and we need access to the bits anyway.



Redistribution

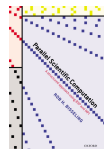


- We redistribute the vector \mathbf{x} from group-cyclic distribution with **cycle** c_0 to **cycle** c_1 , where $c_0 | c_1$ (and hence $c_0 \leq c_1$).
- Optimisation: vector components are sent in packets, not individually.
- BSP model: no difference in cost.
- BSPlib implementation: using packets is more efficient, and gives **optimistic g -values**.



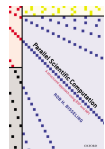
Regular parallel algorithms

- The communication pattern of a **regular parallel algorithm** can be predicted exactly and each processor can determine exactly where every communicated data element goes.
- For a regular algorithm, it is always possible for the user to combine data for the same destination in a block, or **packet**, and communicate the block using **1 put operation**.
- This requires packing at the source processor and unpacking at the destination processor.



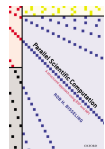
Anything you can do, I can do better

- Song from the musical Annie Get Your Gun, Irving Berlin, 1946.
- The BSP system packs data, but for regular algorithms the user can do better, saving header information that identifies the data.
- This is worthwhile if the communication pattern involves sending many single data words, as happens in the FFT, or many very small data quantities.
- Anything you can send
I can send faster.
I can send anything
Faster than you.
- Not everything you can do, you should do.



How to pack

- Leave this to someone else. Good packers in theory make bad packers in practice.
- If you can leave it up to the BSP system, that's OK too.
- Main question: **which data move to the same processor?**
- Consider x_j and $x_{j'}$ residing on the same processor in the old distribution with cycle c_0 . They are in the same block of size $\frac{nc_0}{p}$ handled by a group of c_0 processors.
- Each block of the old distribution fits entirely in a block of the new distribution, because $c_0 | c_1$.
- Thus, x_j and $x_{j'}$ will automatically be in the same new block of size $\frac{nc_1}{p}$ handled by a group of c_1 processors.



When are x_j and $x_{j'}$ on the same processor?

- Write

$$j = j_2 \frac{c_0 n}{p} + j_1 c_0 + j_0.$$

Because j_2 and j_0 depend only on the processor number, which is the same for j and j' , we can write

$$j' = j_2 \frac{c_0 n}{p} + j'_1 c_0 + j_0.$$

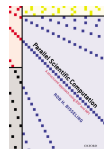
- In the new distribution:

x_j and $x_{j'}$ are on the same processor

$$\iff j \equiv j' \pmod{c_1}$$

$$\iff j_1 c_0 \equiv j'_1 c_0 \pmod{c_1}$$

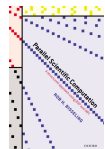
$$\iff j_1 \equiv j'_1 \pmod{\frac{c_1}{c_0}}.$$



Putting one packet

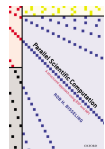
$$j = j_2 \frac{c_0 n}{p} + j_1 c_0 + j_0$$

- The local index of vector component x_j on its processor is $j = j_1$.
- x_j and $x_{j'}$ on the same processor in the new distribution $\iff j_1 \equiv j'_1 \pmod{\frac{c_1}{c_0}} \iff j \equiv j' \pmod{\frac{c_1}{c_0}}$.
- Thus, we can pack components with local indices $j, j + \frac{c_1}{c_0}, j + \frac{2c_1}{c_0}, \dots$, into a temporary array and then put all of these components together into the destination processor as one packet.
- We define **ratio** = c_1/c_0 , which is the stride for packing data.



How not to unpack

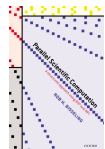
- **Unpacking** is moving data from the location they were put into, to their final location on the same processor.
- If x_j and $x_{j'}$ are two adjacent components in a packet, with **local indices at the source** satisfying $j' = j + \text{ratio}$, then the **global indices** satisfy $j' = j + \frac{c_1}{c_0}c_0 = j + c_1$.
- Thus, the **local indices at the destination** in the group-cyclic distribution with cycle c_1 satisfy $j' = j + 1$.
- We are lucky: if we put the **first component** x_j of the packet **directly** into its final location, and the next component of the packet into the next location, and so on, then all components of the packet immediately reach their final destination.
- Hence, we do not have to unpack!



Redistribution (simplified)

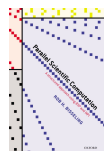
```
void bspredistr(double *x, int n, int p, int s,
               int c0, int c1){ ...
    np= n/p;                ratio= c1/c0;
    size= MAX(np/ratio,1);   npackets= np/size;
    j0= s%c0;                j2= s/c0;

    tmp= vecallocd(2*size);
    for(j=0; j<npackets; j++){
        jglob= j2*c0*np + j*c0 + j0;
        destproc= (jglob/(c1*np))*c1 + jglob%c1;
        destindex= (jglob%(c1*np))/c1;
        for(r=0; r<size; r++){
            tmp[2*r]= x[2*(j+r*ratio)];
            tmp[2*r+1]= x[2*(j+r*ratio)+1];
        }
        bsp_put(destproc,tmp,x,destindex*2*SZDBL,
                size*2*SZDBL);
    } bsp_sync();   vecfreed(tmp);
```



Main function `bspfft` (forward)

```
void bspfft(double *x, int n, int p, int s,  
           double *w0, double *w, double *tw,  
           int *rho_np, int *rho_p){  
    ...  
    np= n/p; k1= k1_init(n,p); rev= TRUE;  
    permute(x,np,rho_np);  
    for(r=0; r<np/k1; r++)  
        ufft(&x[2*r*k1],k1,1,w0);  
    c0= 1; ntw= 0;  
    for (c=k1; c<=p; c *=np){  
        bspredistr(x,n,p,s,c0,c,rev,rho_p);  
        rev= FALSE;  
        twiddle(x,np,1,&tw[2*ntw*np]);  
        ufft(x,np,1,w);  
        c0= c; ntw++;  
    }  
}
```



Summary

- We have optimised the communication in the only communication function the parallel FFT program has, the **redistribution**. This function is crucial for the parallel performance.
- The optimisation is done by **packing** data, which is always possible for **regular algorithms** with a predictable communication pattern.
- Where possible, we have moved computations to initialisation functions, e.g. for the table of weights.
- Because all communication is isolated in one function, the program can easily be ported to another communication library such as MPI.

