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***A Sparse Knapsack Algo-tech-cuit and its Synthesis***

Rumen Andonov, Sanjay Rajopadhye

**N° 2260**

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PROGRAMME 1

Architectures parallèles,  
bases de données,  
réseaux et systèmes distribués



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# A Sparse Knapsack Algo-tech-cuit and its Synthesis

Rumen Andonov\*, Sanjay Rajopadhye\*\*

Programme 1 — Architectures parallèles, bases de données, réseaux  
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Projet API

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**Abstract:** We systematically derive an improved algorithm (called the sparse algorithm) for the general knapsack problem which has better average case performance than the standard (dense) dynamic programming algorithm. The derivation is based on transformation of the standard recurrences into a stream functional programs, and cannot be achieved by the usual space-time mapping techniques because the dependencies are statically unpredictable. Furthermore such a sparse algorithm for the general knapsack problem has not been proposed in the literature, to the best of our knowledge. We also implement the sparse algorithm on a linear asynchronous array with constant size memory on each PE (i.e., a wavefront array processor). The algorithm can run on any number of processors and has optimal time speedup. The second (backtracking) phase of the dynamic programming is also modified so that it can be performed in sparsely and in a memory efficient manner.

*(Résumé : tsvp)*

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# Un algorithme “creux” pour le problème du sac à dos: synthèse et réalisation

**Résumé :** Nous montrons comment dériver systématiquement un algorithme amélioré (appelé *algorithme creux*) pour le problème général du sac-à-dos avec des performances meilleures en moyenne que l’algorithme standard par programmation dynamique. Cette dérivation, qui repose sur une traduction des équations récurrentes standard vers un langage fonctionnel de listes, ne peut faire appel aux techniques classiques de projection spatiale et temporelle car les dépendances ne sont pas connues statiquement. Un tel algorithme creux pour le problème du sac à dos général n’apparaît pas à notre connaissance dans la littérature. Nous implémentons aussi cet algorithme sur un réseau linéaire asynchrone (un réseau à fronts d’ondes) dans lequel chaque processeur élémentaire dispose d’une mémoire de taille fixe. Cet algorithme creux fonctionne avec un nombre quelconque de processeurs et offre un *speedup* optimal. La seconde phase (le retour arrière) de la programmation dynamique est également modifié pour s’exécuter aussi de manière creuse et économe en mémoire.

# 1 Introduction

The knapsack problem is a classic, NP-complete combinatorial optimization problem with many applications [Hu69, MT90], and can be formulated as follows: we are given a knapsack of capacity  $c$ , into which we may put  $m$  types of objects, each object of type  $i$  has a *profit*,  $p_i$ , and a *weight*,  $w_i$ , ( $w_i$ ,  $p_i$ ,  $m$  and  $c$  are all positive integers). Determine  $z_i$ , the number of  $i$ -th type objects to be chosen so as to maximize the total profit without exceeding the capacity, i.e.,

$$\max \left\{ \sum_{i=1}^m p_i z_i : \sum_{i=1}^m w_i z_i \leq c, z_i \geq 0 \text{ integer}, i = 1, 2, \dots, m \right\} \quad (1)$$

If  $z$  is restricted to be a *binary* vector (i.e., each object can be chosen at most once) we have the well known 0/1 case. The knapsack problem can be solved sequentially in  $\Theta(mc)$  time, and this gives it a *pseudo polynomial* complexity [GJ79] since it is polynomial w.r.t. a parameter (which itself, may grow exponentially with input size).

We study dynamic programming [Bel57, Hu69], one of the standard approaches to solving this problem. The algorithm works in two phases, a forward phase for calculating the *profit* of the optimal solution, and then, a backtracking phase that uses this to construct the solution vector,  $z$ . The recurrence equation for the forward phase is given below.

$$f_k(j) = \max(f_{k-1}(j), p_k + f_k(j - w_k)) \quad (2)$$

The equation is defined for all  $0 < j \leq c, 0 < k \leq m$ , and the boundary conditions are,  $f_0(j) = f_k(0) = 0$  and  $f_k(j) = -\infty$  when  $j < 0$ ; we are interested in finding  $f_m(c)$ . The recurrence for the 0/1 knapsack problem is only a slight modification (see the subscript of the second argument):

$$f_k(j) = \max(f_{k-1}(j), p_k + f_{\lfloor \frac{k-1}{2} \rfloor}(j - w_k)) \quad (3)$$

As compared to branch and bound (another classic technique to solve the problem), it is often argued [MT90] that dynamic programming calculates the function  $f_k(j)$  at *all* points and thus *always* pays the worst case price,  $\Theta(mc)$ , whereas the average case performance of branch and bound algorithms is often much better. However, Horowitz and Sahni [HS74] have given a dynamic programming algorithm for the 0/1 problem which has much better average case performance. To date, such an improved algorithm has not been developed for the *general* knapsack problem. In this paper, we systematically derive such an algorithm, and then address its implementation on an application specific processor array. Like the Horowitz-Sahni algorithm, our improved algorithm also uses the fact that the cost function is monotonic—hence one needs to calculate it only at certain *critical points*. The contributions of the paper are summarized as follows:

- An improved algorithm (called the sparse algorithm) for the general knapsack problem which has better average case performance than the standard (dense) dynamic programming algorithm.
- Systematic derivation of the sparse algorithm by transformation of the recurrence of the standard dynamic programming formulation into a stream functional program.
- Modification of the algorithm for bookkeeping information so that the backtracking phase can also be performed sparsely in a very memory efficient manner (because of space limitations, we do not discuss this here. The interested reader is referred to [AR94b], available for anonymous ftp from `ftp.irisa.fr`).
- Implementation of the sparse algorithm on a linear asynchronous array with constant size memory on each PE (i.e., a wavefront array processor [KAGR82]) by proving bounds on the size of local buffer memory in each processor.

The remainder of the paper is organized as follows: Sec 2 gives an intuitive explanation of the Horowitz-Sahni 0/1 knapsack algorithm. Then, Sec 3 presents the formal derivation of the sparse algorithm for the general knapsack problem (because of space limitations we only treat the forward phase). Sec 4 shows how the algorithm can be implemented on a wavefront array processor, where each PE has only finite memory. Finally, Sec 5 gives our conclusions and describes open problems. We conclude this introduction with a brief survey.

## 1.1 Related Work

Many authors have investigated parallel (dense) dynamic programming algorithms for the knapsack problem. Most work has concentrated on the 0/1 problem (for example, [Fer88]), and addresses different models of parallel machines—PRAMs, distributed memory machines, linear arrays, meshes and perfect shuffles. There has also been some work on the general knapsack problem, all dealing with the dense dynamic programming algorithm [AR94a, CCJ90, Ten90]. For this, the best performance is obtained by our systolic algo-tech-cuit [AR94a] (see references therein for a complete discussion).

The basic idea [NU69] of the sparse algorithm is a quarter century old, but seems to have been used only for the 0/1 problem, to the best of our knowledge. Horowitz and Sahni give sequential algorithms for the 0/1 knapsack and related problems using this idea [HS78] (Sec 5.5) [HS74], and Lee et al. implement it on a hypercube using a divide and conquer strategy [LSS88], which takes  $O(mc/q + c^2)$  time on  $q$

processors, and uses  $O(mc)$  storage in the worst case\*. Chen et al. present a pipelined linear array which uses  $\Theta(mc)$  storage,  $\Theta(c)$  on each of  $m$  processors [CJ92, CM92]. They claim that the backtracking takes  $O(m)$  time, but do not give an algorithm to substantiate this claim.

We mention that our model of computation is weaker—an asynchronous ring of  $q$  processors, where each processor has only a constant memory (of size  $\alpha$ ), and we achieve the optimal worst-case performance, namely  $\Theta(\frac{mc}{q})$ . Unlike some of the theoretical work, we do not attempt to bring the running time down to polynomial at the expense of an exponential number of processors (even with the rapid advances in VLSI technology, we feel that this is too expensive).

## 2 Intuitive Explanation of the Sparse Algorithm

We now present an intuitive explanation of the Horowitz-Sahni sparse algorithm for the 0/1 knapsack problem. In the recurrence (3),  $f_k(j)$  represents the optimal profit that can be achieved using only the objects  $1 \dots k$ , and with a capacity of  $j$ . If we add more capacity (i.e., as  $j$  increases), we can only *improve* the total profit, and we thus have the following observation.

**Remark 1** *The function  $f_k(j)$  is monotonically increasing in  $j$ .*

Now, any monotonic function has a “stepwise” shape (see Fig 1), and hence, one can determine its value for any  $j$ , if we know its values at the start of each step. It may therefore be represented completely by a set,  $S_f$  of *pairs* (called critical points)  $[x, F(x)]$ , giving the coordinates where it changes, and the new value.

The sparse algorithm is based on two key observations:

**Remark 2** *If  $F(x)$  is a monotonic function, then, for positive integers,  $p$  and  $w$ , the function  $p + F(x - w)$  is also monotonic.*

In fact, it is simply the function,  $F(x)$  shifted to the right by  $w$  and up by  $p$  (see Fig 2). As a result, it can also be represented by a set,  $S'_f$  of pairs.  $S'_f$  is constructed from  $S_f$  by simply adding the pair  $[w, p]$  to each element and by including  $[w, p]$  as an additional element in the set:

$$S'_f = \{[w, p]\} \cup \{[x + w, F(x) + p] \mid [x, F(x)] \in S_f\}$$

**Remark 3** *If  $F(x)$  and  $G(x)$  are two monotonic functions, then the function  $H(x) = \max(F(x), G(x))$  is also monotonic.*

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\*Lin and Storer point out that this could be worse than the sequential algorithm [LS91]. Note however, that the average behavior is expected to be better because of sparsity.



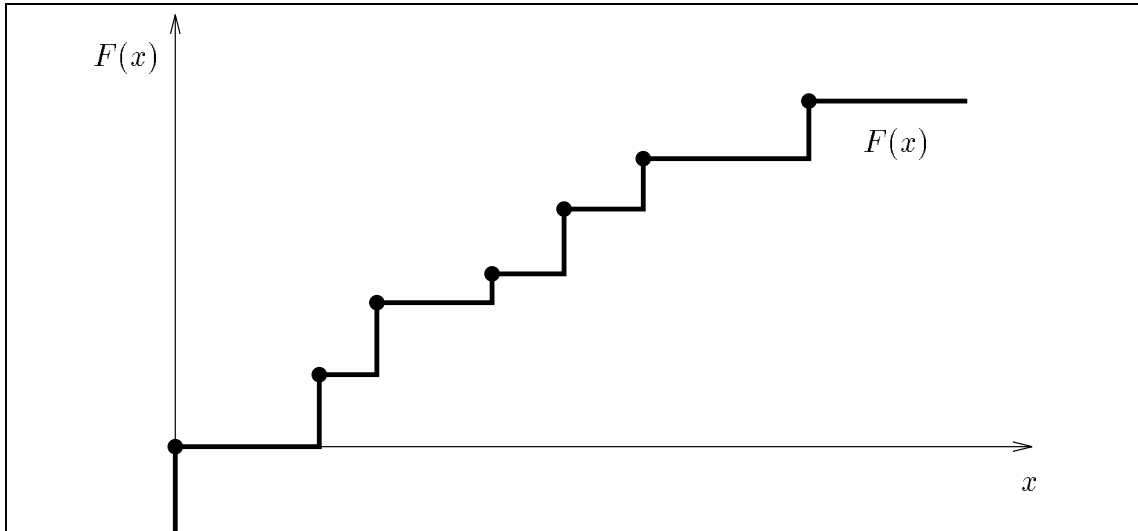


Figure 1: A monotonic function  $F(j)$  can be described by a set of “critical points”.

Fig 3 illustrates this, where  $F(x)$  and  $G(x)$  are shown in thin lines (their critical points are circles and squares, respectively). The max of the two functions is simply the “outer envelope” as shown by thick line.  $H(x)$  is also monotonic, and its critical points are either the circles or squares (i.e., only critical points of one of the two

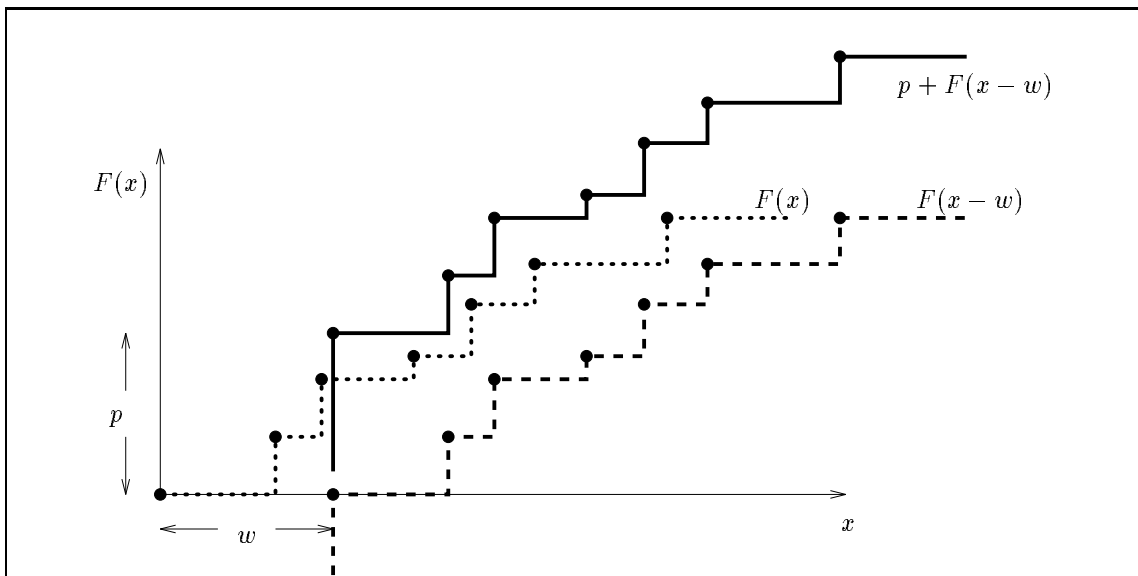
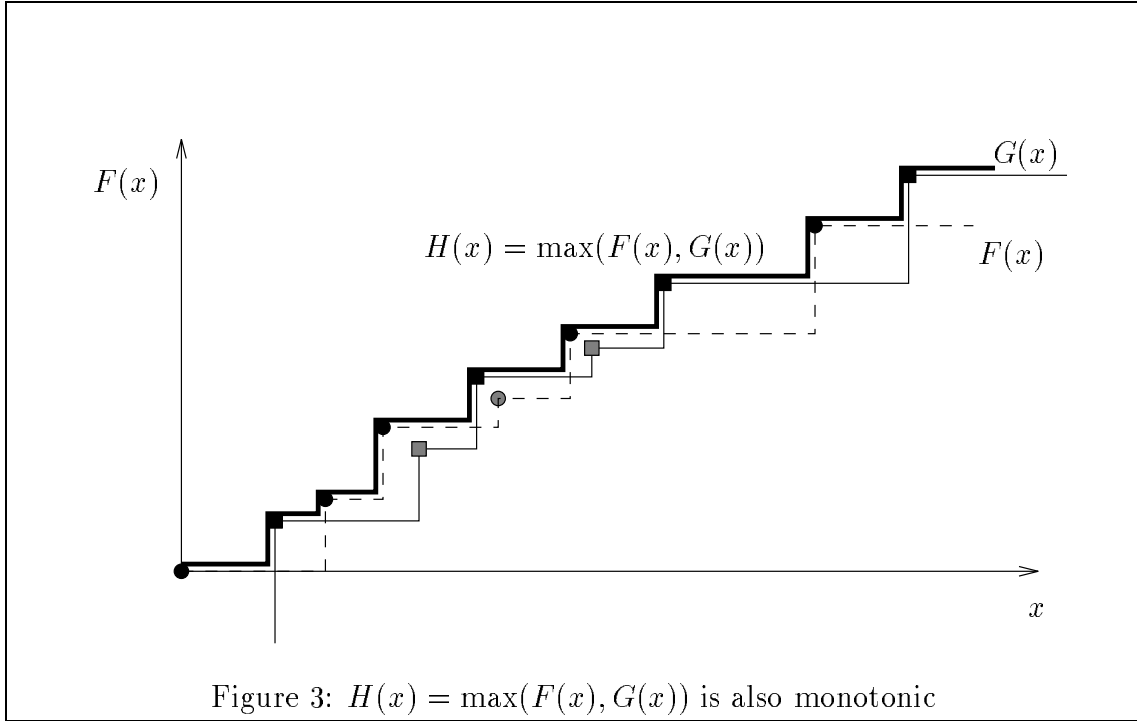


Figure 2:  $F(j)$  is monotonic (shown dotted); so are  $F(x - w)$ —shifted to the right (dashed) as well as  $p + F(x - w)$ —shifted right *and* up (solid).



functions). Note also that some of the critical points (of  $F(x)$  as well as  $G(x)$ ) may not appear (shown in gray) in  $H(x)$ .

Hence we have the algorithm given below. Our explanation has been based on that of Horowitz and Sahni (see [HS78] (Sec. 5.5) for additional details).

1.  $S_1 = \{[w_1, p_1]\}$
2. for  $k = 2 \dots m$ , construct  $S_k$  as follows:
  - (a) Let  $S' = S_{k-1} \cup \{[w_k, p_k]\} \cup \{[x + w_k, f + p_k] \mid [x, f] \in S_{k-1} \wedge x + w_k \leq c\}$  (this is a superset of the critical points of  $S_k$ ).
  - (b) Construct  $S_k$  by removing all critical points  $[x, f] \in S'$  that are *dominated* by another, i.e., if  $\exists [x', f'] \in S' \mid x' \leq x$  and  $f' \geq f$ .
3. Now  $f_m(c)$  can be easily obtained from  $S_m$ .

### 3 Formal Derivation of the Sparse Algorithm

Although the basic idea behind the above algorithm is over a quarter century old [NU69], it has not been applied to the general knapsack problem, to the best of our knowledge. It is clear that Remarks 1–3 are still valid, as illustrated in Tables 1 and 2.

$k \backslash y$	1	2	3	4	5	6	7	8	9	10	11	12	13	14
1	0	0	0	0	7	7	7	7	7	14	14	14	14	14
2	0	0	0	8	8	8	8	16	16	16	16	24	24	24
3	0	0	0	8	8	9	9	16	16	17	17	24	24	25
4	0	0	0	8	8	9	9	16	16	24	24	24	24	32

Table 1: Values of  $f_k(j)$  for a simple general knapsack problem:  $m = 4$ ,  $c = 14$ ,  $w_i = 5, 4, 6, 10$ ;  $p_i = 7, 8, 9, 24$  showing the monotonicity.

$k$	
1	[5,7] [10,14]
2	[4,8] [8,16] [12,24]
3	[4,8] [6,9] [8,16] [10,17] [12,24] [14,25]
4	[4,8] [6,9] [8,16] [10,24] [14,32]

Table 2: The pairs used to represent the example of Table 1.

However, the presence of  $f_k$  on the right hand side of (2) precludes a naive extension, since now,  $S_k$  depends (recursively) on itself. Therefore we need to develop an algorithm that constructs it *incrementally* using the partially computed  $S_k$ . For this reason, we represent the function as a stream of pairs rather than a set. Furthermore, we maintain two additional invariants in our representation: the pairs are sorted in increasing order of weights,  $j$ , (and hence also the associated,  $f$ 's); and the  $j$ 's are bounded by  $c$ , the knapsack capacity.

Let, for  $k = 1 \dots m$ ,  $\mathbf{S}(k)$  be the sequences that represent  $f_k(j)$ . We now systematically derive a functional program that operates on sequences of pairs and calculates  $\mathbf{S}(k)$  from  $\mathbf{S}(k-1)$ . We use the following notation for writing such programs: sequences are delimited by braces (so,  $\{\}$  denotes the empty sequence);  $\mathbf{x} \hat{\mathbf{S}}$  denotes a sequence whose *head* is the element  $\mathbf{x}$ , and whose *tail* (i.e., the rest of the sequence) is the sequence,  $\mathbf{S}$ ; the elements of our sequences in this paper are pairs of integers,  $[j, f]$  (we use brackets for constructing pairs). Function application uses

parentheses (eg,  $\text{Merge}(A,B)$  is the result of applying the function  $\text{Merge}$  to the two arguments  $A$  and  $B$ . We define functions equationally, using separate equations for different patterns of the inputs; for example, the following specifies the behavior of the  $\text{Merge}$  function, when either of its inputs is the empty sequence:

$$\begin{aligned}\text{Merge}(\{\}, S) &= S \\ \text{Merge}(S, \{\}) &= S\end{aligned}$$

Our derivation is based on the following series of lemmata, which follow directly from Remarks 1–3, the discussion in the previous section, and the invariants of our representation.

**Lemma 1** *If  $f_k(j)$  is represented by the sequence  $S(\mathbf{k})$ , then,  $p_k + f_k(j - w_k)$  is represented by  $\text{AddTest}([\mathbf{wk}, \mathbf{pk}], [0,0]^{\wedge}S(\mathbf{k}), c)$  where*

$$\begin{aligned}\text{AddTest}([\mathbf{w}, \mathbf{p}], \{\}, c) &= \{\} \\ \text{AddTest}([\mathbf{w}, \mathbf{p}], [j, f]^{\wedge}S, c) &= \text{if } j + \mathbf{w} > c : \{\} \\ &\quad \text{else} : [j + \mathbf{w}, f + \mathbf{p}]^{\wedge} \text{AddTest}([\mathbf{w}, \mathbf{p}], S, c)\end{aligned}$$

**Lemma 2** *If  $F(x)$  and  $G(x)$  are monotonic functions represented by  $S1$  and  $S2$ , respectively, then  $H(x) = \max(F(x), G(x))$  is represented by a subsequence of  $\text{Merge}(S1, S2)$ , defined as follows:*

$$\begin{aligned}\text{Merge}(\{\}, S) &= S \\ \text{Merge}(S, \{\}) &= S \\ \text{Merge}([j_1, f_1]^{\wedge}S1, [j_2, f_2]^{\wedge}S2) &= \\ \quad \text{if } j_1 < j_2 &: [j_1, f_1]^{\wedge} \text{Merge}(S1, [j_2, f_2]^{\wedge}S2) \\ \quad \text{else if } j_2 < j_1 &: [j_2, f_2]^{\wedge} \text{Merge}([j_1, f_1]^{\wedge}S1, S2) \\ \quad \text{else if } j_1 = j_2 &: [j_1, \max(f_1, f_2)]^{\wedge} \text{Merge}(S1, S2)\end{aligned}$$

Note that in the last clause above, the head of both sequences is consumed and a single output element is produced. This ensures that the output produced by the  $\text{Merge}$  is *strictly* increasing in its  $j$  values, a fact that will be useful later. Thus, the function  $\text{Merge}$  achieves the equivalent of Step 2(a) of the sparse 0/1 algorithm. Now, all that is left is to remove the dominated elements. Recall that a pair  $[j', f']$  dominates  $[j, f]$  iff  $j' \leq j \wedge f' \geq f$ , i.e., it has smaller (or equal) weight but larger (or equal) profit. Because the output of  $\text{Merge}$  is in order of increasing weights, the dominator (if any) of an element must be one of its *predecessors in the sequence*. Hence we have the following.

**Theorem 1** *The following program computes  $S_k$  from  $S_{k-1}$ .*

$$S(\mathbf{k}) = \text{Filter}(\text{Merge}(S(\mathbf{k}-1), \text{AddTest}([\mathbf{wk}, \mathbf{pk}], [0,0]^{\wedge}S(\mathbf{k}), c)), 0)$$

where

```

Filter({}, limit) = {}
Filter([j,f]^S, limit) = if f > limit : [j,f]^Filter(S, f)
                        else           : Filter(S, limit)

```

Note that  $S(k)$  occurs on the right hand side, so the program defines the sequence recursively. Furthermore, the structure of the program is a standard one that occurs in lazy functional languages [AW76, FW76], and is often used to manipulate infinite sequences (without having to evaluate them). The final solution of the (forward phase of the) general knapsack problem is obtained from  $S(m)$ , with the initialization that  $S(0)=\{\}$ .

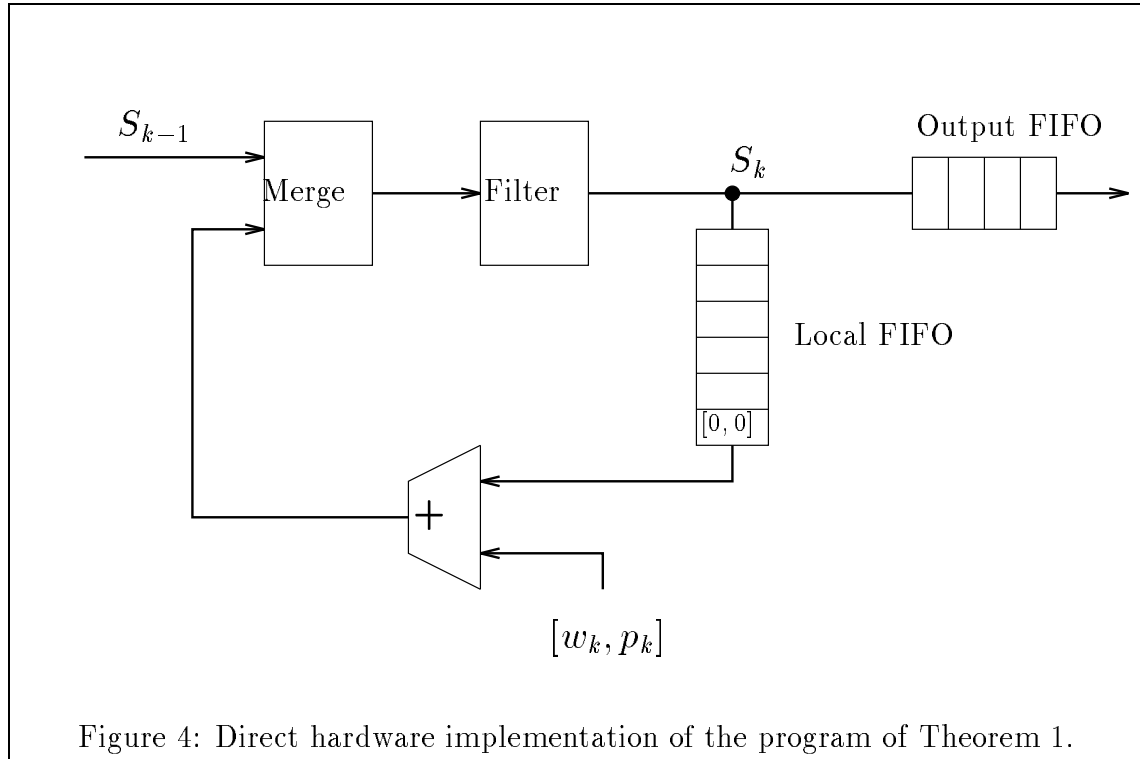
## 4 A Linear Wavefront Array Processor

It is well known [Kah74] that a stream functional program can be implemented as a network of processes, communicating asynchronously through fifo buffers. Such a network can be systematically constructed based only on the *syntactic structure* of the program. For example, the processor that implements our program to compute  $S(k)$  from  $S(k-1)$  is constructed as follows. Our output is the result of applying the function `Filter` (with a scalar parameter, 0) to the output of a `Merge` unit. The latter takes two arguments, one of which is the input sequence, and the other is the output of an `AddTest` unit. The input of this is itself the output sequence  $S(k)$  (with the pair  $[0,0]$  at its head). Hence we need a feedback loop which consists of a fifo buffer, initialized with  $[0,0]$  (see Fig 4).

Looking at the definitions of these three functions, we see that `Filter` can be implemented using a simple “threshold” register which is initialized to 0. If the  $f$  value on the input pair is greater than the threshold, it is passed through (updating the threshold register to  $f$ ), otherwise it is consumed without producing an output (we say that it is *filtered out*). The `Merge` unit is straightforward, and is implemented with two comparators (the second one is used to compare  $f$  values in case the  $j$ 's are equal). `AddTest` is implemented simply by a pair of adders, and a comparator for generating an “end of sequence” tag (not shown in the figure).

Now, to implement the recurrence (2), we need  $m$  such processors, connected in a linear array (if fewer processors are available, the algorithm can be implemented by configuring them as a ring and using multiple passes). Each processor is connected to the next through an *output buffer*, and has a local feedback buffer described above.

It can be shown that such a network of processors is a *partially correct* implementation of the given program—if the program terminates, then the network of processes also terminates and produces the same values. All our arguments in the previous section dealt with partial correctness of the program, and we need to prove that the program of Theorem 1 does terminate. To show this we need to prove one crucial fact, namely that it is free of deadlock (because of the recursive nature of the



program, this is a distinct possibility). We will show this by proving a lower bound on the size of the buffers.

A second important aspect is the memory requirement, i.e., the size of the buffers. For a software implementation on a general purpose machine, this is not a limitation. However, our target is a dedicated VLSI processor, and it becomes crucial to prove that the buffers are bounded from above. The following theorem answers these two problems for the feedback buffer.

**Theorem 2** *At any time (except at the end), the number of elements in the local queue of the  $k$ -th processor is bounded from above by  $w_k$ , and from below by 1.*

**Proof:** Let, at any instant, the local buffer contain elements  $[j_1, f_1], \dots, [j_l, f_l]$ ; the buffer length is thus  $l$ . Also, let the value in the threshold register of the **Filter** unit be  $f'$ , and let  $[j, f]$  be the value at the head of the input sequence. Now, we have the following four cases.

1. **Merge** selects the local value,  $[j_1, f_1]$ , which is (a) filtered out, or (b) passed through by **Filter**.
2. **Merge** selects the input value,  $[j, f]$ , which is (a) filtered out, or (b) passed through by **Filter**.

In case 1.a,  $[j_1, f_1]$  is deleted from the front of the buffer, and the size of the local buffer decreases by 1. In case 1.b,  $[j_1, f_1]$  is deleted from the front of the buffer, and  $[j_1 + w_k, f_1 + p_k]$  is placed at the end, so the buffer size remains  $l$ . In case 2.a also, it remains the same (the input value,  $[j, f]$  is simply deleted). Finally, in case 2.b it increases by 1 ( $[j, f]$  is placed at the end of the buffer, without changing the front). We now consider case 1.a to show a lower bound, and 2.b to show an upper bound.

Now, the fact that the value in the threshold register is  $f'$  means that there is a pair  $[j', f']$ , in the local buffer (which has been passed through some time in the past\*). In case 1.a,  $f_1 + p_k \leq f'$  (by definition of being filtered out), and thus the pair  $[j', f']$  is a *successor* of  $[j_1, f_1]$ . Hence, whenever the buffer size decreases, there is at least one other value in the local buffer that has not yet been consumed.

Now, for case 2.b, we have the following constraints:

$$\begin{aligned} j_1 + w_k &> j \\ j &\geq j_1 + l \end{aligned}$$

The first constraint is because  $[j, f]$  is the winner, while the second is because the output sequence is monotonic and there are  $l - 1$  elements between  $[j_1, f_1]$  and  $[j, f]$ . Hence,  $l < w_k$ . Since the buffer is initialized with the singleton,  $[0, 0]$ , its length is *always* bounded from above by  $w_k$ . ■

Concerning the size of the output buffer, observe that the processors are connected linearly, only through these buffers. This means that the flow of data is unidirectional. Now, if an output buffer is full, the processor is forced to wait. However, we can show by simple induction that its output buffer will eventually become non-full (assuming that the host is continuously consuming data from the output buffer of the last processor). The processor is thus idle only for a bounded number of steps, and there is no possibility of deadlock. The output buffer size cannot affect the correctness of the implementation, but only the speed (latency). In practice the size can be determined by extensive simulation. Our preliminary results indicate that a very small number (two or three registers) is usually sufficient.

## 4.1 Implementation with Fixed Size Buffers

From Theorem 2, we now have a linear asynchronous array whose memory is bounded by  $w_{\max}$ , the largest possible weight, a problem parameter. Because of this, our solution is not completely modular—if  $w_{\max}$  increases, it will be necessary to redesign the

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\*Note that initially,  $f' = 0$  and the local buffer is initialized with  $[0, 0]$

processor. However, we can use the approach that was previously used for deriving a pure systolic array for the dense knapsack problem [AQRW94]. If we have processors with buffer space for only  $\alpha$  pairs, we simply configure  $\lceil w_i/\alpha \rceil$  physical processors to simulate one processor. Since all the  $w_i$  values are known when a problem is loaded, we can also determine (off line) a few bits of configuration information which informs the PE whether it is to participate actively or simply contribute its memory (it can be easily shown that two bits are enough following the lines of [AQRW94]). Such an array with  $q$  processors, configured into a ring, can solve any problem instance in (note that this is the worst case result)  $\left\lceil \frac{1}{q} \sum_{i=1}^m \lceil w_i/\alpha \rceil \right\rceil$  passes. The worst case running time is then  $\Theta(\frac{mc}{q})$ , which is optimal.

## 5 Conclusions

In this paper, we have used some very old ideas to derive a sparse algorithm for the dynamic programming technique for solving the general knapsack problem. The basic idea is the fact that the cost function is monotonic [NU69], and it suffices to calculate it only at certain *critical points*, i.e., points where the function changes value. Because of this, the function can be represented as sequences of pairs. The sequences are, on the average, much shorter than the knapsack capacity, and this is the key to the improvement of the average running time. Horowitz and Sahni have presented extensive empirical data to support this fact for the sparse 0/1 algorithm, and our preliminary simulations confirm that this is also true for the general problem. Also note that since the problem is NP complete, the *worst case* time is still exponential (pathological examples can be easily constructed [Chv80]). An analytic study of the average case complexity is the subject of our ongoing work.

Another contribution of the paper is the systematic derivation of the algorithms by transforming the original recurrence equation into stream functional programs. Our derivation is almost mechanical, and provides a proof of correctness of the derived programs.

Finally, we have shown how the stream functional programs that we derived could be implemented on a wavefront array processor, an asynchronous processor array, where each PE has fixed memory, independent of problem parameters. In such an array, problems with larger parameters sizes are simply solved by adding more processors, or reusing the same array in multiple passes. In order to do this we prove that the lengths of the buffers in architecture is always bounded. This raises an interesting open question. It is well known that lazy stream functional programs [FW76], correspond almost mechanically to networks of processes communicating through fifo channels [Kah74], these are not realistic because the problem of proving bounds on buffer lengths is undecidable [Wad84]. We were successful in proving the



bounds for our example by hand, and it would be interesting to see if the ideas used in the proof could be generalized to certain cases of asynchronous processor arrays.

Finally, it is necessary to analytically determine a good model for the average case behavior of the sparse algorithms and hence the processor arrays. If a closed form for this can be determined, one could try to optimize the expected running time by appropriately choosing the buffer size in each PE. This approach has been successfully used for the dense algo-tech-cuit [AR94a], and it would be interesting to extend it to the sparse case.

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