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Articles

Using Problem Structure in Derivative-free Optimization

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1. Introduction

Derivative-free optimization is the branch of optimization where the minimizer of functions of several variables is sought without any use of the objective's derivatives. Although a number of problems may involve constraints, we will focus in this paper on the unconstrained case, i.e.,

$$\min_{x \in \mathbb{R}^n} f(x), \quad (1)$$

where f is an objective function which maps \mathbb{R}^n into \mathbb{R} and is bounded below, and where we assume that the gradient of f (and, a fortiori, its Hessian) cannot be computed for any x .

The main motivation for studying algorithms for solving this problem is the remarkably high demand from practitioners for such tools. In most of the case known to the author, the calculation of the objective function value $f(x)$ is typically *very expensive*, and its derivatives are not available either because $f(x)$ results from some physical, chemical or econometrical measure, or, more commonly, because it is the result of a possibly very large computer simulation, for which the source code is effectively unavailable. The occurrence of problems of this nature appears to be surprisingly frequent in the industrial world. For instance, we have heard of cases where evaluating $f(x)$ requires the controlled growth of a particular crop or the meeting of an adhoc committee. As can be guessed from this last example, the value of $f(x)$ may in practice be contaminated by noise, but the numerical techniques for handling this latter characteristic are also outside the scope of our presentation.

Several classes of algorithms are known for derivative-free optimization. A first class is that of direct search techniques, which includes the well-known and widely used simplex reflection algorithm of [22] or its modern variants [11, 27], the old Hookes and Jeeves method [20] or the parallel direct search algorithm initiated by Dennis and Torczon [15, 29]. These methods are based on a predefined geometric pattern or grid and use this tool to decide where the objective function should be evaluated. A recent and comprehensive survey of the development in this class is available in [21] and the associated bibliography is accessible at <http://www.cs.wm.edu/~va/research/wilbur.html>. See also the paper by Abramson, Audet and Dennis in this issue. The main advantage of methods in this class is that they do not require smoothness (or even continuity) of the objective function.

A second algorithmic class of interest is that of interpolation/approximation techniques pioneered by Winfield [30, 31] and by Powell in a series of papers starting with [23, 24]. In these methods, a low-order polynomial (linear or quadratic) model of the objective function is constructed and subsequently minimized, typically in a trust-region context (see [6, Chapter 9]). Because they build smooth models, they are appropriate when the objective function is known to be smooth. A common feature of the algorithms of this type is that they construct a basis of the space of suitable polynomials and then derive the model by building the particular linear combination of the basis polynomials that interpolate (or sometimes approximate) known values of $f(x)$. Powell favors a basis formed of Lagrange fundamental polynomials, while Conn *et al.* [7, 8, 10] use Newton fundamental polynomials instead. Both choices have their advantages and drawbacks, which we will not discuss here. The polynomial space of interest is typically defined as the span of a given set of monomials: full quadratic polynomials in the two variables x_1 and x_2 are for instance those spanned by 1, x_1 , x_2 , x_1x_2 , x_1^2 and x_2^2 .

While methods in these two classes have been studied and applied in practice, their use has remained essentially limited to problems involving only a very moderate number of variables: the solution of problems in more than 20 variables is indeed possible in both cases, but is typically very compu-

tationally intensive. In direct search techniques, this cost is caused by the severe growth in the number of grid or pattern points that are necessary to “fill” \mathbb{R}^{20} or spaces of even higher dimensions. A similar difficulty arises in interpolation methods, where a total of $(n+1)(n+2)/2$ known function values are necessary to define a full quadratic model, but it is also compounded with the relatively high complexity in linear algebra due to repetitive minimization/maximization of such quadratic polynomials. One may therefore wonder if there is any hope for the derivative-free solution of problems in higher dimensions. It is the purpose of this paper to indicate that this hope may not be unfounded, at least for problems that exhibits some structure. Section 2 discusses the partially separable structure which will be central here, while Sections 3 and 4 indicate how it can be numerically exploited in direct search and interpolation methods, respectively. We then consider the special case of sparse problems in Section 5. Some conclusions are finally presented in Section 6.

2. Problem structure

Large-scale optimization problems often involve different parts, or blocks. Each block typically has its own (small) set of variables and other (small) sets of variables that link the block with other blocks, resulting, when all blocks are considered together, in a potentially very large minimization problem. One may think, for instance, of a collection of chemical reaction tanks, each with its temperature, pressure or stirring controls, and with its input raw material and output products. Other examples include discretized problems where a given node of the discretization may involve more than a single variable but is only connected to a few neighbouring nodes, or PDE problems with domain decomposition, or multiple-shooting techniques for trajectory/orbit calculations, where each orbital arc only depends on a few descriptive or control variables, with the constraint that the different arcs connect well via a (small) set of common variables. Examples of this type just abound, especially when the size of the problem grows. It is interesting that their structure can very often be captured by the notion of *partial separability*, introduced by Andreas Griewank and the author in [17, 19].

A function $f : \mathbb{R}^n \rightarrow \mathbb{R}$ is said to be *partially separable*, if its value may be expressed, for every x , as

$$f(x) = \sum_{i=1}^q f_i(U_i x), \quad (2)$$

where each *element function* $f_i(x)$ is a function depending on the *internal variables* $U_i x$ with U_i being a $n_i \times n$ matrix ($n_i \ll n$). Very often, the matrix U_i happens to be a row subset of the the identity matrix, and the internal variables are then just a (small) subset of the original problem variables.

Griewank and Toint show in [18] that *every twice-continuously differentiable function with a sparse Hessian is partially separable*, which gives a hint at why such functions are so ubiquitous. It is remarkable that, although this is only an existence result and specifies a particular decomposition (2) only indirectly, we are not aware of any practical problem having a sparse Hessian and whose partially separable decomposition is not explicitly known.

Partially separable functions (and their extension to group-partially separable ones) have been instrumental in the design of several numerical codes, including the LANCELOT package [5]. We now intend to show how they can be exploited in derivative-free optimization.

3. Direct search for partially separable objective functions

For our description of the structure use in direct search methods, we focus on the common case where the matrices U_i are subsets of the identity, which then implies that each element function f_i only depends on a small subset of the problem variables. As in [28], we assume that each f_i is available individually, along with a list of the problem variables on which f_i depends. Two problem variables are then said to interact if at least one element f_i depends on both. Sets of non-interacting variables are useful because the change in f from altering all variables in the set is just the sum of the changes in f arising from altering each variable in the set individually.

Exploiting partial separability is advantageous for direct search methods in two separate ways. It can both reduce the computational cost of each function

value sought, and also provide function values at related points as a by-product. For example, consider a totally separable function

$$f(x) = \sum_{i=1}^n f_i(x_i)$$

where $x = (x_1, \dots, x_n)^T$, and where the value of each element f_i is known at a point x . Assume, as is typical in pattern search methods, that we wish to explore f in the vicinity of x by calculating f at each point of the form $\{x \pm he_i\}_{i=1}^n$, where h is a positive constant and e_i is the i^{th} column of the identity matrix. Calculating f at each point requires the evaluation of one f_i , and so the total cost of the $2n$ function values at the points $\{x \pm he_i\}_{i=1}^n$ amounts to *two* function evaluations of the complete function f . However, any set of changes to f from steps of the form $\pm he_i$ are independent of one another provided no two such steps alter the same variable. Hence as a by-product we obtain the function values at all other points of the form $x + h \sum_{i=1}^n \eta_i e_i$ where each η_i is either $-1, 0$, or 1 . Thus we obtain f at $3^n - 1$ new points for the total cost of *two* function evaluations.

When f is only partially separable, the gains are not as dramatic as in the totally separable case, but they still can be very substantial. For example, let f be of the form

$$f(x) = f_1(x_n, x_1, x_2) + \sum_{i=2}^{n-1} f_i(x_{i-1}, x_i, x_{i+1}) + f_n(x_{n-1}, x_n, x_1). \quad (3)$$

Then, given each $f_i(x)$, calculating f at $\{x \pm he_i\}_{i=1}^n$ costs the equivalent of six complete function evaluations. When n is divisible by 3, an inductive argument shows that the number of other function values obtained as by-products is $7^{(n/3)} - 2n - 1$.

A third example shows that the two advantages of partial separability are distinct. Let $f = f_3(x_1, x_2) + f_2(x_1, x_3) + f_1(x_2, x_3)$. Calculating f at all points of the form $x \pm he_i$, $i = 1, 2, 3$, costs four function evaluations, given that each $f_i(x)$, $i = 1, \dots, 3$ is known in advance. No further function values are generated as by-products.

We may now exploit these features in constructing a pattern search method along the lines described in [11], where minimization is conducted on a sequence

of successively finer nested grids (each grid is a subset of its predecessor), aligned with the coordinate directions. The m^{th} grid $\mathcal{G}^{(m)}$ is of the form

$$\mathcal{G}^{(m)} = \left\{ x^{(0)} + 2^{1-m} \sum_{i=1}^n \eta_i e_i \mid \eta_i \in \mathbb{Z} \right\},$$

where $x^{(0)}$ is the initial point and \mathbb{Z} is the set of signed integers. Loosely speaking, each grid is obtained by taking its predecessor and ‘filling in’ all points half-way between each pair of existing grid points.

For each grid, the algorithm generates a finite subsequence of iterates on the grid, with monotonically decreasing objective function values. This subsequence is terminated when no lower grid point can be found around the current iterate (the final iterate is then called a grid local minimizer) and the algorithm then proceeds onto the next grid in the sequence.

The key idea here is to exploit partial separability in constructing the minimizing subsequences. This is done as follows. Before the first iteration, the algorithm starts by grouping the problem variables into subsets indexed by V_1, \dots, V_r such that all variables whose index are V_p appear in exactly the same element functions f_i 's. This means that all these variables are equivalent in terms of which other variables they interact with. (In example (3), $V_p = \{p\}$ for $p = 1, \dots, r = n$.) Each V_p then determines a subspace

$$S_p = \text{span}\{e_j\}_{j \in V_p}.$$

Thus, identifying these subspaces with their generating variables, we have that some S_p interact and some (hopefully most) are *non-interacting*. (In example (3) again, the only interactions are between S_{p-1}, S_p and S_{p+1} and between S_1 and S_n .) We next build a positive basis B_p for each S_p , that is a set of vectors such that any $v \in S_p$ can be written as a positive linear combination of the vectors of B_p . For V_p , this basis can be chosen, for instance, as

$$B_p = \{e_j\}_{j \in V_p} \cup \left\{ - \sum_{j \in V_p} e_j \right\},$$

in which case it is also minimal (see [12, 14] for further developments on positive basis). Minimization on a given grid $\mathcal{G}^{(m)}$ is then achieved by repeating the following steps.

1. For each $p \in \{1, \dots, r\}$, we first calculate the objective function reduction

$$\Delta_p = \min_{v_j \in B_p} \sum_{i=1}^q \left[f_i(x^{(k)} + v_j) - f_i(x^{(k)}) \right],$$

and denote by s_p the argument of the minimum. Because of our definition of the subspaces S_p , only a small fraction of the elements $f_i(x^{(k)} + v_j)$ must be computed for every v_j .

2. We next choose the best objective function reductions from a set of non-interacting subspaces, i.e., we select an index set $I \subseteq \{1, \dots, r\}$ such that

$$\Delta_I = \sum_{p \in I} \Delta_p$$

is minimal and the subspaces $\{S_p\}_{p \in I}$ are non-interacting.

3. If $\Delta_I < 0$, we finally define

$$x^{(k+1)} = x^{(k)} + \sum_{p \in I} s_p,$$

(note that $f(x^{(k+1)}) = f(x^{(k)}) + \Delta_I$), and increment k .

This is repeated until a grid local minimum is located ($\Delta_I \geq 0$). The procedure is then stopped if the desired accuracy is reached, or, if this not the case, the grid is refined and a new grid minimization started.

Of course, this description of the algorithm remains schematic, and we refer the interested reader to [28] for further details and algorithmic variants. In particular, this reference describes a technique for computing the index sets V_1, \dots, V_r from the partially separable structure of the objective function. We also note that it is not crucial to solve the combinatorial problem of Step 2 exactly: we may, for instance, use a simple greedy algorithm to identify a suitable index set I .

Does this work in practice? Table 1 shows the performance in terms of complete function evaluations of the method just described (under the heading ‘‘PS’’, for partial separability), compared to the same algorithm without the exploitation of problems structure (under the heading ‘‘no st.’’). The comparison is made on two well-known test problems from the CUTer test set [16].

n	LMINSURF		BROYDN3D	
	PS	no st.	PS	no st.
9	215	501	343	1241
16	483	3724	334	3605
25	484	10557	364	8087
36	890	19796	379	16503
49	1002	—	363	24531
64	1149	—	362	—
81	1413	—	389	—
100	1634	—	362	—
121	2045	—	362	—
144	2120	—	392	—
169	2689	—	361	—
196	3233	—	361	—
5625	79511	—	535	—

Table 1: Number of function evaluations required to minimize the linear minimum surface (LMINSURF) and Broyden tridiagonal (BROYDN3D) functions in various dimensions (from [28]).

The conclusion is very clear: using the partially separable structure in direct search methods makes their application to relatively large-scale problems possible, while the unstructured approach rapidly reaches its limits in size. It is also interesting to note that the methodology adapts in an obvious way if derivatives of some but not all element functions are available, or if one is ready, for some or all p , to compute Δ_p by minimizing in the complete subspace S_p instead of only considering the grid points defined by the positive basis B_p .

4. Interpolation methods for partially separable problems

We now turn to interpolation techniques and focus on trust-region methods. At each iteration of this type of methods, (typically quadratic) model of the objective function is constructed, which interpolates a set of known functions values. In other words, one builds the quadratic $m^{(k)}$ such that

$$m^{(k)}(y) = f(y) \quad \text{for all } y \in Y^{(k)},$$

where $Y^{(k)}$ is the *interpolation set* at iteration k , a set of points at which values of the objective are known. As indicated above, the number of points in $Y^{(k)}$ must be equal to $(n+1)(n+2)/2$ for estimating a fully quadratic $m^{(k)}$. This model is then minimized

within the trust-region, and the resulting trial step accepted or rejected, depending on whether or not the achieved decrease in f sufficiently matches the predicted decrease in the model. The size of the trust-region is then reduced if the match is poor, or possibly enlarged if it is satisfactory. This broad algorithmic outline of course hides a number of practical issues that are critical to good numerical performance. The most important is that the geometry of the sets $Y^{(k)}$ (i.e., the repartition the interpolation points y in \mathbb{R}^n) must satisfy a condition called *poisedness*. If we consider a two-dimensional problem, for instance, it is rather intuitive that the points in $Y^{(k)}$ should not all lie on a straight line. This poisedness condition can be formalized and tested [8, 9, 26] in various ways, but the key observation is that the same set of points cannot be used for ever as the algorithm proceeds, or even only modified to include the objective values at the new iterates. Figure 1 illustrates the potentially disastrous effect on the model of a bad geometry (bottom), compared to a situation at a previous iteration where the geometry is adequately poised (top).

As a consequence, it is necessary to improve the geometry of $Y^{(k)}$ at some iterations, typically by computing the objective function's value at new points chosen to ensure sufficient poisedness.

How can we adapt this technique if we now assume that we know a partially separable decomposition (2) of the objective? (We no longer assume that the matrices U_i are row subsets of the identity.) The answer, which is fully developed in [4], is based on a fairly obvious observation: *instead of building $m^{(k)}$, a model of f in the neighbourhood of an iterate $x^{(k)}$, we may now build a collection of models $\{m_i^{(k)}\}_{i=1}^q$, where each $m_i^{(k)}$ now models f_i in the same neighbourhood.* Estimating a structured quadratic model of the form

$$m^{(k)}(x^{(k)} + s) = \sum_{i=1}^q m_i^{(k)}(x^{(k)} + s)$$

then requires at most $(n_{\max} + 1)(n_{\max} + 2)/2$ complete function evaluations, where we have defined $n_{\max} = \max_{i=1, \dots, q} n_i$. Since $n_{\max} \ll n$, this is typically order(s) of magnitude less than what is required for estimating an unstructured model. Moreover, n_{\max} is often independent of n .

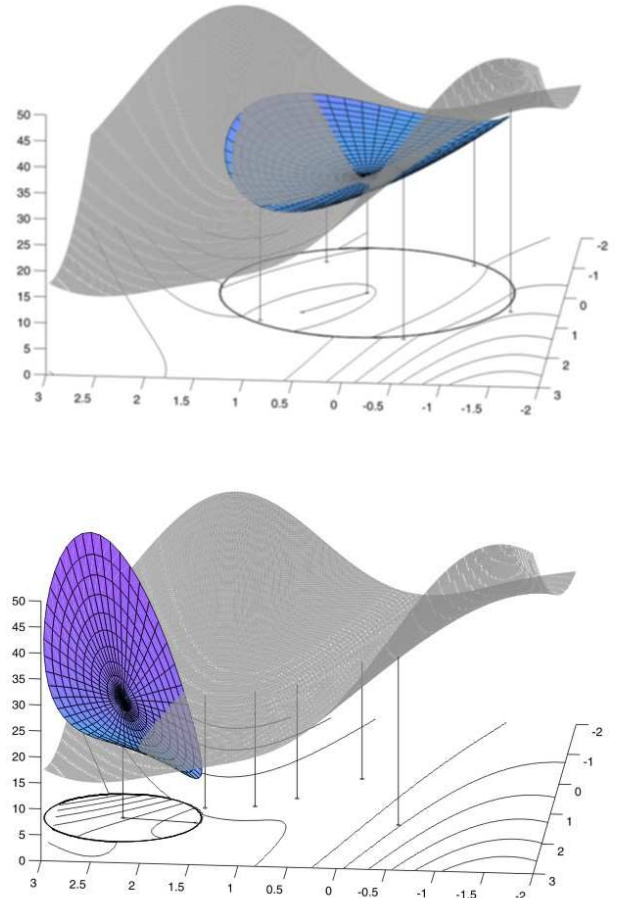


Figure 1: How the evolution of the interpolation set can lead to a bad model when its geometry deteriorates (the interpolation points are materialized by vertical bars from the surface to the plane, the model is quadratic and shown inside the trust-region)

However, we now have to manage q interpolation sets $Y_i^{(k)}$ ($i = 1, \dots, q$) over the iterations, instead of a single one. We then have to consider two possible cases: either it is possible for the user to evaluate a single element function $f_i(x)$ independently of the others (as we have assumed above), or the objective function must always be evaluated as a whole (i.e., only the collection of values $\{f_i(x)\}_{i=1}^q$ can be computed for a given x). In the second case, a straightforward implementation of the algorithm could be very expensive in terms of function evaluations if one blindly applies a geometry improvement procedure to each $Y_i^{(k)}$. Indeed, computing a vector that

improves the geometry of the i -th interpolation set yields a vector having only n_i “useful” components, and it is not clear which values should be assigned to the remaining $n - n_i$ components. The strategy in this case is to group the necessary individual function evaluations by applying the CPR procedure of Curtis, Powell and Reid [13] for estimating sparse Jacobian matrices to the $n \times q$ occurrence matrix of variables x_j into elements f_i . (Other techniques involving graph colouring in the spirit of [1] are also possible). This typically results in substantial savings in terms of function evaluations.

As in the previous section, we do not elaborate more on the algorithmic details and variants (and refer to [4] for this), but rather illustrate the potential benefits of the approach on a few examples. These benefits are already visible for medium-size problems, as is shown in Table 2 where the unstructured algorithm (UDFO) is compared with its version using partially separable structure (PSDFO) on two CUTEr problems. The “I” version of PSDFO refers to the case where each element function f_i is accessible, and the “G” version to the case where the complete collection of $\{f_i(x)\}_{i=1}^q$ must be evaluated together for every x .

Problem	n	UDFO	PSDFO(I)	PSDFO(G)
ARWHEAD	10	311	30	54
	15	790	30	56
	20	1268	35	161
	25	1478	37	198
BDQRTIC	10	519	348	358
	15	1014	345	382
	20	1610	509	596
	25	2615	360	542

Table 2: Comparison of the number of function evaluations required by DFO solvers for solving medium-size instances of problems ARWHEAD and BDQRTIC (from [4]).

Again, the use of the partially separable structure brings clear benefits, and these are also more pronounced when the values of the elements can be computed individually (case “I”). Table 3 indicates that these benefits extend to higher dimensions, as expected.

n	GENHUMPS		BROYDN3D	
	PSDFO(I)	PSDFO(G)	PSDFO(I)	PSDFO(G)
10	114	168	53	70
20	200	345	56	84
50	202	357	68	114
100	249	433	153	342
200	253	436	123	278

Table 3: The number of function evaluations required by PSDFO for solving problems GENHUMPS and BROYDN3D in various dimensions (from [4]).

5. Interpolation methods for sparse problems

We conclude our overview of the use of structure in derivative-free optimization by examining interpolation methods in the less favourable case where structure is present but access to the individual values of the element functions $f_i(x)$ impossible. This may happen, for instance, when the objective function value results from a complicated simulation involving a discretized partial differential equation (such as a complicated fluid calculation). In such cases, we often know that a partially separable decomposition of the form (2) exists (resulting, in our example, from the discretization topology), but we are only given the final value of $f(x)$, without its decomposition in its element function values (note the difference with the “G” case in the previous section). This implies that we typically know that the Hessian of f , $H(x)$, is sparse for every x , and know its structure. Can we exploit this (more limited) information?

As above, the idea is to construct a quadratic model $m^{(k)}$ that reflects the problem structure as much as possible: we thus need to construct a model whose Hessian has the same sparsity structure as that of the objective function. Interestingly, as noted in [2, 3], this is remarkably easy. Indeed, a given sparsity pattern is equivalent to a selection of a subset of the monomials generating the quadratic polynomials: if the (i, j) -th entry of $H(x)$ (and thus the (j, i) -th one) is known to be zero for all values of x , this simply indicates that f can be modelled by a restricted quadratic polynomial that does not involve the monomial $x_i x_j$. Thus the models $m^{(k)}$ may now be viewed as a linear combination of $1+n+n_H$ monomials, where n_H is the number of nonzeros in the

lower triangular part of $H(x)$. This last number is often a small multiple of n , in which case the size of the interpolation set $Y^{(k)}$ is linear rather than quadratic in n . The cost of evaluating a sparse quadratic model is thus also very attractive, although typically larger than that of a partially separable model (often independent of n , as we have noted in the previous section).

These observations have been exploited in [2, 3], but also in [26] where Powell describes his successful UOBSQA code. The efficiency of this technique is attested by the results of Table 4, where the number of objective evaluations required for convergence is reported for UDFO (as in Table 2) and UOBSQA.

Problem	Dimension	UDFO	UOBSQA	UOBDQA
ARWHEAD	$n = 10$	311	118	105
	$n = 15$	790	170	164
	$n = 20$	1268	225	260
	$n = 25$	1478	296	277
BDQRTIC	$n = 10$	519	350	288
	$n = 15$	1014	594	385
	$n = 20$	1610	855	534
	$n = 25$	2615	1016	705

Table 4: Comparison of the number of function evaluations required by UDFO, UOBSQA and UOBDQA solvers for medium-size instances of problems ARWHEAD and BDQRTIC (from [4]).

Finally, one may argue that there is in fact no need that the sparsity structure of the model's Hessian really reflects that of $H(x)$. It is indeed possible to simply *impose an a priori sparsity structure* in order to reduce the size of the interpolation set, even if no sparsity information is available for $H(x)$. This idea was suggested in [2] and implemented, in its extreme form where the model's Hessian is assumed to be diagonal, in the UOBDQA code by Powell [25]. The excellent results obtained with this technique are also apparent in Table 4, even if one might guess that it will be mostly effective on problems whose Hessian is diagonally dominant. Potential extensions of this idea include the exploration of techniques (inspired for instance from automatic learning) which adaptively tailor the imposed sparsity pattern to the effective behaviour of the objective function.

6. Conclusions and perspectives

Given the discussion above, one may clearly conclude that *yes, the derivative-free optimization algorithms can exploit problem structure to (sometimes dramatically) improve their efficiency*. Moreover, the arguments presented also indicate that a richer structural information typically results in more substantial efficiency gains when using interpolation methods: the number of objective function evaluations necessary to estimate a quadratic model indeed ranges from quadratic in n (no structure) via linear in n (when exploiting sparsity) to independent of n (when using partial separability).

It is clear that the development of structure-aware derivative-free optimization methods and packages is only starting, and much remains to be done. We think, in particular, of extensions of the ideas discussed here to the constrained case, and their applications to neighbouring research areas, such as domain decomposition (an ongoing project explores the use of partial separability in this context) and others. Developments in these directions combine both the more abstract aspects of algorithm design and theory with the very practical nature of a subject in high industrial demand. There is no doubt that they therefore constitute valuable research challenges.

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