# PCP Characterizations of NP: Towards a Polynomially-Small Error-Probability

Irit Dinur<sup>\*</sup>

Eldar Fischer<sup>†</sup> Guy Kindler<sup>‡</sup> Ran Raz<sup>§</sup> Shmuel Safra<sup>‡</sup>

January 25, 2010

#### Abstract

This paper strengthens the low-error PCP characterization of NP, coming closer to the upper limit of the BGLR conjecture. Consider the task of verifying a witness for the membership of a given input in an NP language, using a constant number of accesses. We show that it is possible to achieve an error probability exponentially small in the number of bits accessed, where the number of bits in each access is as high as  $\log^{\beta} n$ , for any constant  $\beta < 1$ . The BGLR conjecture asserts the same for a constant  $\beta$  where  $\beta \leq 1$ .

Our results are in fact stronger, implying that the Gap-Quadratic-Solvability problem with a constant number of variables in each equation is NP-hard. That is, given a system of n quadratic-equations over a field  $\mathcal{F}$  of size up to  $2^{\log^{\beta} n}$ , where each equation depends on a constant number of variables, it is NP-hard to distinguish between the case where there is a common solution to all of the equations, and the case where any assignment satisfies at most a  $\frac{2}{|\mathcal{F}|}$  fraction of them.

At the same time, our proof presents a *direct* construction of a low-degree-test whose error-probability is exponentially small in the

<sup>\*</sup> School of Mathematics, Institute for Advanced Study, Princeton, NJ. .

<sup>&</sup>lt;sup>†</sup> The Faculty of Computer Science, Technion, Israel.

<sup>&</sup>lt;sup>‡</sup> School of Mathematical Sciences, Tel Aviv University, Israel.

<sup>&</sup>lt;sup>§</sup> Weizmann Institute of Science, Israel.

number of bits accessed. Such a result was previously known only relying on recursive applications of the entire PCP theorem.

# 1 Introduction

Cook-Levin's characterization of NP implies that every  $L \in NP$  is reducible to 3-SAT. The reduction from L to 3-SAT is a polynomial-time algorithm that receives an input string I, and produces a set  $\Psi$  of boolean functions (called local-tests), each depending on a constant number of variables.  $\Psi$  represents the membership of I in L, in the sense that there exists an assignment satisfying all local-tests if and only if  $I \in L$ .

A PCP characterization of NP differs from Cook-Levin's characterization in regards to what is guaranteed in the case where the input is not in L: In Cook's characterization, one can only be sure that the reduction will produce a system that cannot be entirely satisfied. To characterize NP in terms of PCP, it must be guaranteed that the reduction algorithm produces a system  $\Psi$  such that no assignment can satisfy even a small fraction  $\epsilon$  of its local-tests.

In both cases, a satisfying assignment to  $\Psi$  can be viewed as a witness for *I*'s membership in *L* (and hence  $\Psi$  can be viewed as a membershipverification system). In a PCP framework, however, this witness can be efficiently verified by randomly picking a local-test of  $\Psi$  and verifying that it holds (hence the term PCP – Probabilistic Checking of Proofs). In this case, the *error probability* parameter,  $\epsilon$ , of the PCP, bounds the probability of accepting *I* even though  $I \notin L$ . Other parameters of  $\Psi$ , such as the variable range and the number of variables accessed by each local-test, are also part of the PCP characterization.

For many applications of PCP, the characterization of NP with a constant error-probability and variables of a constant range [AS92, ALM<sup>+</sup>92] suffices. In order to prove NP-hardness of other problems, however, sub-constant error-probability has turned out to be essential. For example, [LY94] and [BGLR93] were able to prove that approximating SET-COVER to within logarithmic factors is *almost* NP-hard, using the constant error-probability PCP characterization of NP. To improve this result to strict NP-hardness, [BGLR93] had suggested the "sliding scale" conjecture.

The sliding scale conjecture states that it is possible to keep the number of variables accessed by each local-reader constant, and to make the variables' range non-constant, obtaining an error probability polynomially small in the size of the variable-range. In other words, it is possible to achieve a membership verification system for any NP-language where each local-test accesses a constant number of 'words' (variables), and where the error-probability is exponentially small in the 'word-length' (number of bits in each variable).

One cannot expect the error-probability to be less than polynomially small in the size of the variables' range, since a random assignment will satisfy any satisfiable local-test with such a probability (recall that each test depends on a constant number of variables). Hence the sliding scale conjecture is optimal in the sense of error-probability.

According to the conjecture, the variables' range may be increased up to a size polynomial in the length of the original input (note that each local-test can be given as a truth-table). Reaching larger range-size is unlikely since the error-probability would then become less than  $1/|\Psi|$ . In the case where the input is not in L, this implies that no local-test succeeds, so the problem of deciding whether the input is in L reduces to that of deciding whether any of the local-tests is satisfiable.

The sliding scale conjecture was shown to hold for a sizable portion of the applicable range-size in [RS97], where a PCP characterization of NP was shown that achieves error-probability polynomially small in the size of the variable range for a variable range of size up to  $2^{\log^{\beta} n}$ , where  $\beta < 1$  is a certain positive constant (see also [AS97]).

## **Our Main Results**

In this paper, we prove the sliding scale conjecture for variable range sizes of up to  $2^{\log^{\beta} n}$  where  $\beta$  is *any* constant smaller than one (as opposed to "*some* constant" achieved by [RS97]), thus coming closer to proving the sliding scale conjecture for the full applicable range.

In fact our result is somewhat stronger, proving the conjecture for the aforementioned range using proof verification systems of a specific structure. In these systems the local-tests have the form of quadratic-equations instead of being general boolean functions, with the variables' range representing a finite field. This result implies that for a quadratic equation-system of n equations over a field  $\mathcal{F}$  (with  $|\mathcal{F}| \approx 2^{\log^{\beta} n}$  for any fixed constant  $\beta < 1$ ), where each equation depends on a constant number of variables, it is NP-hard to decide whether there exists a common solution to all equations, or whether any assignment to the variables satisfies no more than a  $\frac{2}{|\mathcal{F}|}$  fraction

of them.

One of the main tools used to obtain the above result, which is interesting in its own right, is that of LDF-readers, a version of what is known in the literature as a low-degree test (see [RS97, AS97]). A direct construction of an LDF-reader is shown herein, that achieves an exponentially-small errorprobability with respect to the number of bits it accesses. Such LDF-readers could previously be attained only by recursive applications of the entire PCP theorem.

# **Related Results**

We note that there is no known PCP characterization of NP, where the size of the variable-range is polynomial in the size of the membership-verification system (or equivalently, the length of each variable is logarithmic in it), and the error probability is exponentially small in the number of accessed bits. This is true even when allowing a super-polynomial time reduction. The repetition lemma of [Raz98] shows that by two accesses to  $\Theta(\log n)$  bits, the error-probability can be made polynomially small in n, where n is the size of the original input, while the size of the generated system is  $n^{\log n}$ . Similarly, the multi-linear extension of [BFL91] yields a system with a  $\frac{1}{n}$ error-probability, whose size is  $n^{\log n}$ . In fact, in any known reduction there is always a factor of at least  $\log^{\epsilon} n$  in the exponent that separates the errorprobability from the size of the generated instance.

Achieving an error-probability polynomially small in the size of the generated instance is an important open problem. Such a characterization of NP would improve hardness results for several problems. For example, approximating the 'Monotone-Minimum-Satisfying-Assignment' problem (which is closely related to approximating the length of propositional proofs [ABMP98]) has been shown to be NP-hard in [DS98] via a reduction from PCP, such that the hardness of approximation ratio is preserved. Hence a polynomially small error-probability PCP characterization of NP would immediately imply that it is NP-hard to approximate the length of propositional proofs to within an  $n^{\epsilon}$  factor for some constant  $\epsilon > 0$ .

[RS97] managed to keep the exponential relation between the number of bits accessed and the error-probability, thus showing the sliding scale conjecture true for a variable range of size up to  $2^{\log^{\beta} n}$  for *some* constant  $\beta < 1$ . For larger  $\beta$  (any constant  $\beta < 1$ ) [RS97] showed a system whose error probability is  $2^{-\log^{\beta} n}$ , yet without the exponential relation between the number of accessed bits and the error-probability, since the number of bits accessed was  $O(\log^{\beta} n \cdot \text{poly} \log \log n)$ . This factor of poly  $\log \log n$  is significant when viewing, for example, the result in terms of Gap-Quadratic-Solvability. The result of [RS97], if it were to be translated to Gap-Quadratic-Solvability terms, would at best give an equation system with each equation depending on  $O(\text{poly} \log \log n)$  variables. In comparison, our result translates to a quadratic equation-system with the same error-probability, but where every equation depends on a *constant* number of variables, namely  $\Theta(\frac{1}{(1-\theta)^2})$ .

### Techniques

We use the general framework of [AS92, ALM<sup>+</sup>92, RS97] for our proof. However, instead of the generalized form of the composition paradigm utilized in previous PCP proofs, we use a more concrete representation. Our result could have been obtained using the previous structure, but this representation simplifies our proof, and some of its techniques may be of independent interest.

In [HPS93], it was shown that given a system of quadratic equations over a finite field, it is NP-hard to distinguish between the case that the system can be completely satisfied, and the case that not even a small fraction of the equations can be satisfied by a single assignment. The crucial difference between this and our main result is that in the [HPS93] reduction each equation depends on almost all the variables in the system, while our main result claims the same for the case where the equations are restricted to having a constant number of variables each.

Our proof begins with a system  $\Psi$  of quadratic equations as in [HPS93], and reduces it to a system  $\Psi'$  of quadratic equations with a constant number of variables in each. The key property of our proof is that throughout the reduction we use systems of equations over the same field  $\mathcal{F}$ , the field over which  $\Psi$  is defined. The field structure is utilized through various steps of composition, thus enabling us to cross the barrier that limits the proof technique of [RS97].

To simplify the exposition, the reduction partitions the variables of  $\Psi'$ into subsets called domains. In each such domain a mapping is defined, associating each variable to a point in a linear space of the form  $\mathcal{F}^d$  over  $\mathcal{F}$ . An assignment to these variables can thus be regarded as a function over the linear space. The reduction has two main steps. At first, it transforms  $\Psi$  into a system  $\Psi_{sc}$  where the number of variables in each equation is constant. This is accomplished by an iterative application of the sum-check technique from [BFL91]. The system  $\Psi_{sc}$  has the required properties only if the assignment to the variables in each domain, when viewed as a function, is a low-degree polynomial. In order to get rid of this restriction, the reduction then generates LDF-readers and plugs them into the equations of  $\Psi_{sc}$ , thereby obtaining the final system  $\Psi'$ .

**LDF-readers.** LDF-readers are used to obtain evaluations of polynomial functions of low-degree that are represented by a set of variables, by accessing only a very small part of their representation. An LDF-reader should either reject or return values that are consistent with some low-degree polynomial, even if the assignment to the representation variables is not totally consistent with the representation of one polynomial. The probability that, given an incorrect representation, the LDF-reader does not reject but still the returned evaluations are not consistent with a low-degree polynomial, is its error probability. For a more accurate definition of an LDF-reader, the reader is referred to Section 2.

An LDF-reader of sub-constant error-probability seems necessary in order to attain PCP characterizations of NP with sub-constant error-probability. The plane-vs.-plane LDF-reader introduced by [RS97], where a polynomial is represented by its restriction to planes, achieves a sub-constant probability. The previously used line-vs.-point LDF-reader was shown by [AS97] to have a small error-probability as well. However, the error probability of these LDF-readers is not exponentially small in the number of bits they access.

It seems to be difficult to achieve error-probability smaller than polynomial in the number of accessed bits, using a direct LDF-reader comparing subspaces (lines, planes, etc.) for consistency. This is since many bits are required to represent the restriction of a polynomial to a subspace. One way to attain exponentially small error-probability from these LDF-readers is by utilizing the composition technique, applying the entire PCP theorem to them. Our proof, in contrast, makes this recursion concrete, utilizing an explicit representation of low-degree polynomial functions that yields LDF-readers with an exponentially small error probability. The composition-recursion LDF-reader. Our LDF-reader uses a representation of low-degree polynomials as follows. We begin with a representation where a multi-dimensional polynomial is represented by all of its point evaluations, and also by its restriction to certain constant dimensional subspaces. We use a new *power-substitution* technique to then replace each constant dimensional restriction of the polynomial by a multi-dimensional polynomial of a much smaller degree. This is done, roughly, by replacing monomials of high degree with new variables. The polynomials whose degree was reduced are then represented by their point evaluations and their restriction to constant dimensional subspaces, and the process is repeated.

After a constant number of such iterations we obtain polynomials of linear degree over constant dimensional spaces. Each of these polynomials is then represented by a constant number of variables that range over the field  $\mathcal{F}$ . Hence to obtain evaluations our LDF-reader is not required to completely read a low-degree polynomial over some subspace – instead it only accesses a constant number of variables that range over  $\mathcal{F}$ .

#### Organization of the paper

Our main result and the main definitions required for its proof are stated in Section 2. The proof of the main result, based on lemmas that are proven in the following sections, appears in Section 3. The construction of the LDFreader that is utilized in the proof of the main result appears in Section 4. In particular, the power-substitution technique, used in the construction of the LDF-reader to represent polynomials using other polynomials with more variables but with considerably smaller degrees, is described in Subsection 4.4. Finally, Section 5 describes the recursive application of the sum-check (and other) techniques, which are used in the reduction to obtain from the original system  $\Psi$  a system  $\Psi_{sc}$  with a constant number of variables in each equation.

# 2 Preliminaries

In this section we describe the basic ideas and definitions utilized in the proof of our main result.

### Gap-Quadratic-Solvability

The Gap-Quadratic-Solvability problem is that of determining whether all the equations in a given system of quadratic-equations can be simultaneously satisfied, or whether only a small fraction of the equations can be satisfied. Viewing the quadratic equations as local-tests of a PCP system, showing this problem to be NP-hard yields a PCP characterization of NP.

**Definition 1** (Gap-Quadratic-Solvability). The Gap-Quadratic-Solvability problem with parameters D,  $\sigma$  and  $\epsilon$  (which may be, implicitly, functions of the system size n), is denoted by gap- $QS[D, \sigma, \epsilon]$ . An instance of the problem is a fields  $\mathcal{F}$  of size<sup>\*</sup>  $\sigma$ , and a set of n quadratic-equations over  $\mathcal{F}$ , where each equation has at most D variables (D is called the dependency parameter). The problem is to distinguish between the following two cases:

- Yes. There is an assignment to the variables that satisfies all of the equations.
- No. Every assignment to the variables satisfies at most an  $\epsilon$  fraction of the equations  $\epsilon$  is called the error parameter.

An instance which falls under one of the above criteria is said to have the gap property. Any outcome is acceptable for instances that do not have the gap property.

Our main theorem shows NP-hardness of gap-QS with a constant dependency parameter, for a field of size  $\sigma \approx 2^{c \log^{\beta} n}$  and an error parameter  $\epsilon = \frac{2}{\sigma}$ , where  $\beta < 1$  is any constant smaller than 1 and c > 0 is some constant.

Note that the requirement for a  $\frac{2}{\sigma}$  error is almost optimal, since it is easy to satisfy a  $\frac{1}{\sigma}$  fraction of the satisfiable equations of any system by a random assignment. On the other hand, from hardness for any error  $\epsilon = \frac{1}{\sigma^{\Omega(1)}}$ which is polynomially small in the size  $\sigma$  of the field, one can obtain hardness for a  $\frac{2}{\sigma}$  error by a simple amplification technique. We therefore abbreviate gap-QS[ $D, \sigma$ ] for the gap-QS problem where  $\epsilon$  is fixed to be  $\frac{2}{\sigma}$ . Note that this error probability is polynomially small in the size of the field, and therefore exponentially small in the length, measured in bits, of each variable.

<sup>\*</sup>In fact there can be at most one field of any given cardinality, however we would like to be able to look at gap-QS problems where the size can vary in some range. In that case it makes sense to request that the actual field be given as part of the input.

**Theorem 1** (main theorem). For every constant  $\beta < 1$  there exists a constants  $c_2 > c_1 > 0$  such that gap- $QS[O(1), \sigma]$  is NP-hard for  $\sigma$  in the range  $2^{c_1 \log^{\beta} n} \leq \sigma \leq 2^{c_2 \log^{\beta} n}$ , where n is the number of equations in the system.

We actually prove Theorem 1 via a many-to-one reduction. Informally speaking, this means that gap-QS[ $O(1), \sigma$ ] is proven to be NP-complete.

Gap-QS[ $n, \sigma$ ], where the number of variables in each equation is not bounded, is proven to be NP-hard in [HPS93] for any  $\sigma$  that is polynomially bounded in n, using simple linear codes:

**Theorem 2** ([HPS93]). Let  $\sigma(n)$  be any positive poly-time computable function, and suppose that for some constant  $\gamma > 0$ ,  $\sigma(n) = O(n^{\gamma})$ . Then there exist constants  $d_2 > d_1 > 0$  such that  $Gap-QS[n,\sigma]$  is NP-hard for  $\sigma$  in the range  $(\sigma(n))^{d_1} \leq \sigma \leq (\sigma(n))^{d_2}$ .

This theorem is proven by a relatively simple reduction from the Cook-Levin characterization of NP, so the entire difference between this characterization of NP and the PCP characterization boils down to the constant bound on the number of variables that each equation accesses.

Theorem 1 is proven by showing a reduction algorithm, taking as input a system  $\Psi$  of n quadratic-equations and producing a new system  $\Psi'$  where the number of variables in each equation is bounded by a constant. The number of variables in each equation is reduced while roughly preserving the fraction of satisfiable equations. Specifically, if  $\Psi$  is completely satisfiable then  $\Psi'$  is completely satisfiable as well; and if no more than a  $\frac{2}{|\mathcal{F}|}$  fraction of the equations of  $\Psi$  can be satisfied then the same occurs for  $\Psi'$ .

### 2.1 LDFs and Domains

Let us set a notation for polynomial functions of low degree – an object used extensively in this paper.

**Definition 2** (LDF - low degree function). An [r, d]-LDF is a polynomial function from  $\mathcal{F}^d$  to  $\mathcal{F}$ , of total-degree at most r.

The variables of the system  $\Psi'$  that is generated by our reduction all range over  $\mathcal{F}$  (since  $\Psi'$  is a system of quadratic equations over  $\mathcal{F}$ ). However, for the exposition of the reduction algorithm and for the proof of correctness, it is useful to consider certain subsets of these variables as separate *domains*. The variables in each such domain are associated with the points of a vector field  $\mathcal{F}^d$  over  $\mathcal{F}$ , and an assignment to the variables in the domain can thus be viewed as a function  $f: \mathcal{F}^d \to \mathcal{F}$ , as explained in the following definition. Throughout our reduction, the domains are always disjoint, so in the final system  $\Psi'$  and also in all intermediate constructions, no variable can belong to more than one domain at the same time.

**Definition 3** (domain). A domain F is a set of variables ranging over  $\mathcal{F}$ , that has one variable F[x] for every point  $x \in \mathcal{F}^d$ , where d = d(F) is called the dimension of the domain. F is said to be assigned a function  $f : \mathcal{F}^d \to \mathcal{F}$  if for every x, the variable F[x] is assigned f(x).

Two more parameters are associated with each domain in addition to the dimension – the lower-degree, denoted s(F), and the upper-degree r(F)(r(F) will always be larger than s(F)).

Let us stress that the lower-degree and upper-degree parameters of domains, as well as the whole partition of variables into domains, are only figments of our proof and construction. In the final system  $\Psi'$  the domainstructure is discarded, and we are left with variables that range over  $\mathcal{F}$ . These variables can be assigned every value in  $\mathcal{F}$ , and therefore a domain Fcontaining these variables can be assigned any function  $f: \mathcal{F}^{d(F)} \to \mathcal{F}$ .

However, in the proof we give special consideration to assignments of domains which correspond to LDFs. In particular, it will be shown that if the system  $\Psi'$  generated by the reduction is satisfiable, there is a satisfying assignment where every domain F is assigned an s(F)-degree LDF. In case  $\Psi$  cannot be more than  $\frac{2}{|\mathcal{F}|}$  satisfiable, it should be proven that no assignment can satisfy more than  $\frac{2}{|\mathcal{F}|}$  of the equations of  $\Psi'$ . However, we prove later that it suffices to only show this for assignments where each domain F is assigned an r(F)-degree LDF.

**Definition 4** (assignment of a domain). The assignment f of a domain F is said to be good, if f is an [s(F), d(F)]-LDF, and it is said to be feasible if f is an [r(F), d(F)]-LDF. An assignment for a set of variables containing one or more domains is said to be good (feasible) if the assignment to each domain is good (feasible).

The reduction which transforms  $\Psi$  into  $\Psi'$  goes through an intermediate system  $\Psi_1$  where the number of variables in each equation is constant, but

which does not yet have the desired properties. In particular,  $\Psi_1$  might be completely satisfiable even when there exists no assignment satisfying more than a  $\frac{2}{|\mathcal{F}|}$  fraction of the equations of  $\Psi$ . However,  $\Psi_1$  behaves much better if we restrict the set of assignments considered: On one hand if  $\Psi$  is completely satisfiable then not only  $\Psi_1$  is completely satisfiable, but there exists a good satisfying assignment for it. On the other hand, if  $\Psi$  is no more than  $\frac{2}{|\mathcal{F}|}$ satisfiable, then there is no good or even feasible assignment for  $\Psi_1$  satisfying more than a  $\frac{2}{|\mathcal{F}|}$  fraction of its equations.

### 2.2 LDF-Readers

To transform  $\Psi_1$  into the final system  $\Psi'$ , we should prevent it from being satisfiable by assignments where domains are not assigned LDFs. In fact what we manage is a bit weaker (although it suffices). Consider an equation  $\psi \in \Psi_1$  that has the variables  $F[x_1], \ldots, F[x_k]$  in a domain F. To prevent it from being satisfiable by unwanted assignments we use a mechanism called an *LDF-reader*, which is plugged into  $\psi$  in place of these variables. The LDFreader ensures that  $\psi$  either reads evaluations at  $(x_1, \ldots, x_k)$  of an LDF (even if the assignment to F is not feasible) or it is not satisfied.

An LDF-reader evaluating the tuple of points  $(x_1, \ldots, x_k)$  in a domain F has two parts – the *representation*, and the set of *local-readers* which produce the evaluations.

**The representation.** The representation is a set V that contains F and maybe other variables and domains. The variables in V, including those associated with domains in it, are called *representation variables*. The LDF-reader uses the representation variables to produce evaluations. Every good assignment f for F must be extensible to a good assignment for all the variables in V, called the *encoding-assignment* of f. When given the encoding-assignment of a good LDF f, the LDF-reader will always return the evaluations of f.

#### The local-readers

The evaluations at  $(x_1, \ldots, x_k)$  are produced by a set of local-readers, where each local-reader accesses only a constant number of representation variables

– this property is essential since the local-readers are plugged into  $\Psi_1$  to produce  $\Psi'$  and the number of variables in each equation must remain constant. Each local-reader may either produce evaluations, or it may reject if it finds that the assignment is not an encoding-assignment.

**Local-tests and evaluators.** Each local-reader is a pair containing a *local-test* – a conjunction of linear equations over representation variables, and a tuple of k evaluators. Each evaluator is a linear-combination of representation variables. A local-reader is said to *accept* an assignment for the representation variables if the local-test is satisfied by it, and otherwise it is said to *reject* it. For an assignment  $\mathcal{A}$  which is an encoding-assignment of a good LDF f, it is required that all local-readers accept, and also that the *i*'th evaluator in each local-reader evaluates to  $f(x_i)$ .

In case the representation variables are not given a correct encodingassignment, we would like the local-readers to always reject. This is not possible to achieve, however, with local-readers that access a constant number of representation variables, not even if we allow a small fraction of the localreaders to falsely accept. It is in fact not even possible to ensure that localreaders which do not reject return evaluations of a single LDF. What we can achieve (and turns out to be enough), is that apart from a small fraction, the local-readers either reject or return evaluations of one of a short list of LDFs. This is the list of LDFs which are *permissible* with respect to the assignment of F.

**Definition 5** (permissible assignment). An [r(F), d(F)]-LDF f is said to be  $\rho$ -permissible with respect to an assignment of F if for at least a  $\rho$ -fraction of the points x, F[x] is assigned f(x).

We show later that for a wide range of permissibility parameters  $\rho$ , the list of permissible LDFs is bounded by  $O(\rho^{-1})$ . Since the list is only determined by the assignment to F and is independent of the rest of the representation variables, it will be the same for all LDF-readers evaluating tuples in F. This means that all equations that have variables in F will read evaluations that are consistent with one of the LDFs on the relatively short list.

We now give the formal definition of the parameters of an LDF-reader.

**Definition 6** ( $(\rho, \epsilon)$ -LDF-reader). Let  $\mathcal{R}$  be an LDF-reader evaluating a tuple  $(x_1, \ldots, x_k)$  in a domain F, and fix an assignment to its representation-

variables. A local-reader L is said to be  $\rho$ -erroneous if it accepts, and there exists no  $\rho$ -permissible LDF f (with respect to the assignment of F) such that the *i*'th evaluator evaluates to  $f(x_i)$  for all *i*.

 $\mathcal{R}$  is said to be a  $(\rho, \epsilon)$ -LDF-Reader, if for any assignment to the representationvariables, the fraction of  $\rho$ -erroneous local-readers is at most  $\epsilon$ .

# 3 Proof of the Main Theorem

In this section we prove Theorem 1. We show for any constant  $\beta < 1$ , a polynomial time reduction from the problem Gap-QS[ $n, \sigma$ ] with  $\sigma$  in the range  $(\sigma(n))^{d_1} \leq \sigma \leq (\sigma(n))^{d_2}$  and where  $d_1$  and  $d_2$  are as in Theorem 2, to the problem Gap-QS[ $O(1), \sigma$ ] with  $\sigma$  in the range  $2^{c_1 \log^{\beta} n} \leq \sigma \leq 2^{c_2 \log^{\beta} n}$ , where  $c_2 > c_1 > 0$  are some appropriate constants.

The reduction starts with a given system  $\Psi$  of n quadratic equations over a field  $\mathcal{F}$  of size  $2^{d_1 \log^{\beta} n} \leq |\mathcal{F}| \leq 2^{d_1 \log^{\beta} n}$ , with up to n variables in each equation. It then generates, in time polynomial in n, a system  $\Psi'$ over the same field with at most a constant number of variables in each equation. Since the reduction is in polynomial time (and since the number of equations in not decreased by the reduction), m is polynomially equivalent to n, and therefore the size of the field satisfies  $2^{c_1 \log^{\beta} m} \leq |\mathcal{F}| \leq 2^{c_2 \log^{\beta} m}$  for appropriate constants  $c_2 > c_1 > 0$ , as required by Theorem 1.

 $\Psi'$  will have the *completeness* property, that is if the given system  $\Psi$  can be completely satisfied then  $\Psi'$  will be completely satisfiable as well; and the *soundness* property – if  $\Psi$  is no more than  $\frac{2}{|\mathcal{F}|}$ -satisfiable (namely no assignment can satisfy more than a  $\frac{2}{|\mathcal{F}|}$  fraction of its equations), then  $\Psi'$  is at most  $\frac{2}{|\mathcal{F}|}$ -satisfiable as well.

The reduction begins by transforming  $\Psi$  into a system  $\Psi_{sc}$  of quadraticequations where the number of variables in each equation is bounded by a constant, and that has the desired properties only if the assignments for its variables are restricted to being feasible. The transformation of  $\Psi$  into  $\Psi_{sc}$  is done by a *sum-check* algorithm, whose properties are stated in the following Lemma, proven in Section 5.

**Lemma 3.1** (sum-check). There exists a polynomial-time algorithm as follows. It takes as input a system  $\Psi$  of n quadratic equations over a field  $\mathcal{F}$ ,  $|\mathcal{F}| = 2^{\log^{\beta} n}$ , where there are up to n variables in each equation. Given  $\Psi$ , the algorithm generates a system  $\Psi_{sc}$  of quadratic-equations over  $\mathcal{F}$  where each equation has a constant number of variables (some of which in domains), and that has the following properties:

- Completeness: If  $\Psi$  is completely satisfiable then  $\Psi_{sc}$  is completely satisfiable by a good assignment.
- Soundness: If  $\Psi$  is no more than  $\frac{2}{|\mathcal{F}|}$ -satisfiable then  $\Psi_{sc}$  cannot be more than  $\frac{2}{|\mathcal{F}|}$ -satisfied by a feasible assignment.

Moreover, all the domains of  $\Psi_{sc}$  have the same dimension  $d = \Theta(\log^{1-\beta} n)$ , lower-degree s, and upper-degree r, where  $s \leq |\mathcal{F}|^{c_1}$  and  $r \geq |\mathcal{F}|^{c_2}$  for some global constants  $c_1 < c_2 < 1$ .

To transform  $\Psi_{sc}$  into  $\Psi'$ , the reduction generates LDF-readers and plugs them into  $\Psi_{sc}$  as follows. For each equation  $\psi$  of  $\Psi_{sc}$ , that has the variables  $F(x_1), \ldots, F(x_k)$  of a domain F, it generates an LDF-reader evaluating  $(x_1, \ldots, x_k)$  in F. To plug the LDF-reader into  $\psi$ , many copies of  $\psi$  are made, and one of the local-readers is plugged into each copy.

The local-tests are added in conjunction with each copy, and hence a system of *conjunctions* is formed. In addition, some of the gap is lost when the LDF-readers are plugged in – if  $\Psi$  is no more than  $\frac{2}{|\mathcal{F}|}$ -satisfiable, the fraction of satisfiable conjunctions in the system obtained from  $\Psi_{sc}$  might be somewhat higher. A simple amplification technique is hence applied to the system of conjunctions to avoid that, and then each conjunction is replaced by equations, obtaining  $\Psi'$ .

### 3.1 Generating the LDF-Readers

To generate LDF-readers we use a *constructor* algorithm, as defined below.

**Definition 7** (constructor). A constructor is an algorithm that takes as input a domain F and a k-tuple  $(x_1, \ldots, x_k)$  of points in  $\mathcal{F}^{d(F)}$ , where k is a constant<sup>†</sup>. It generates an LDF-reader evaluating  $(x_1, \ldots, x_k)$  in F, in time polynomial in  $|\mathcal{F}|^{d(F)}$ . The number of variables appearing in each local-reader must be bounded by a constant, and so should be the number of equations in

<sup>&</sup>lt;sup>†</sup>By saying that k is constant, we mean that other parameters and properties of the constructor may depend arbitrarily on k. In addition, any parameter of the constructor that depends only on k is considered constant as well

the local-test of each local-reader. In addition, the number of local-readers must only depend on the parameters of F.

Our reduction uses the *Composition-Recursion constructor*, whose properties are stated in the next lemma, to generate LDF-readers. Note that the Composition-Recursion can only be applied to domains with restricted dimension and degree parameters. These restrictions are needed to

The reduction uses a constructor that generates *Composition-Recursion* LDF-readers. The existence of this constructor and the properties of the LDF-readers it generates are stated in the next lemma, which is proven in Section 4.

**Lemma 3.2** (Composition-Recursion constructor). There exists a global constant  $c_g$ ,  $0 < c_g \leq 1/2$ , such that for every  $c_1 < c_2 < 1$  and  $\beta < 1$  the following holds. There exist a constant c > 0, and an LDF-Reader constructor for domains of dimension  $d = \Theta(\log^{1-\beta} n)$ , lower-degree  $s \leq |\mathcal{F}|^{c_1}$ , and upper-degree  $r \geq |\mathcal{F}|^{c_2}$  (the algorithm runs independently of s and r). The LDF-readers generated by the algorithm are  $(\rho, O(\rho^c))$ -LDF-readers, for all  $\rho$ 's which satisfy  $\rho > (r/|\mathcal{F}|)^{c_g} d$ .

Before the LDF-readers are actually generated, we make some small technical alterations to  $\Psi_{sc}$  as follows.

**Uniformization.** The number of variables in each equation of  $\Psi_{sc}$  is bounded by some constant k. This implies that an equation in  $\Psi_{sc}$  may have variables from up to k distinct domains, and that the number of variables it has from each domain is bounded by k. Before generating LDF-readers, let us assume for simplicity that each equation of  $\Psi_{sc}$  has variables from *exactly* k distinct domains, and that it has exactly k variables from each domain. This requires the reduction to add arbitrary variables to the equations, multiplied by zero coefficients.

**LDF-Reader generation.** After the uniformization, the reduction generates the LDF-readers as described above – For each equation  $\psi$  of  $\Psi_{sc}$ , that has the variables  $F(x_1), \ldots, F(x_k)$  in a domain F, it generates an LDF-reader evaluating  $(x_1, \ldots, x_k)$  in F (this takes polynomial time in the size of  $\Psi_{sc}$ ). Note that since all domains in  $\Psi_{sc}$  have the same parameters (dimension d, lower-degree s and upper-degree r, as stated in Lemma 3.1), the number of local-readers in each LDF-reader is the same as well. The representation variables of the LDF-readers are added to the variables of the system, and the local-readers are plugged into the equations of  $\Psi_{sc}$  as described below.

# 3.2 Plugging LDF-Readers In

For each equation  $\psi \in \Psi_{sc}$  there are now k associated LDF-readers – one for each domain it has variables from. The first step in plugging the LDFreaders into  $\Psi_{sc}$  is to replace each such equation  $\psi$  by its representation set  $\mathcal{E}_{\psi}$ , containing conjunctions of quadratic equations that are obtained by plugging local-readers into  $\psi$ .  $\mathcal{E}_{\psi}$  represents  $\psi$  in the sense that an assignment satisfying a large enough fraction of the conjunctions in  $\mathcal{E}_{\psi}$  implies a satisfying assignment for  $\psi$ , as shown in the proof of Claim 3.3 below.

**Generating**  $\mathcal{E}_{\psi}$ . Let  $\psi \in \Psi_{sc}$  be an equation that has variables from the domains  $F_1, \ldots, F_k$ . For each j, let us denote the variables of  $\psi$  in  $F_j$  by  $F_j[x_1^j], \ldots, F_j[x_k^j]$ .  $\psi$  is therefore associated with k LDF-readers  $\mathcal{R}_1, \ldots, \mathcal{R}_k$ , where  $\mathcal{R}_j$  evaluates the tuple  $(x_1^j, \ldots, x_k^j)$  in  $F_j$ . The reduction generates one conjunction in  $\mathcal{E}_{\psi}$  for each choice of k local-readers  $L_1, \ldots, L_k$ , where  $L_j \in \mathcal{R}_j$ . The first equation in each such conjunction, denoted  $\psi'$ , is the quadratic equation obtained from  $\psi$  by replacing each variable of the form  $F_j[x_i^j]$  with the *i*'th evaluator of  $L_j$  (it evaluates  $x_i^j$  in  $F_j$ ).  $\psi'$  is then put in conjunction with the local-tests of the local-readers  $L_1, \ldots, L_k$ .

The system  $\Psi_2$ . Note that the number of conjunctions in  $\mathcal{E}_{\psi}$  is the same for every  $\psi \in \Psi_{sc}$  – it is  $|\mathcal{R}|^k$ , where  $|\mathcal{R}|$  denotes the number of local-readers in each of the LDF-readers we have generated. We set  $\Psi_2$  to be the union of all the sets  $\mathcal{E}_{\psi}$ . Since the number of variables in each local-reader is constant, the number of variables in each conjunction of  $\Psi_2$  is bounded by a constant as well. The system of conjunctions  $\Psi_2$  obviously retains the completeness property of  $\Psi_{sc}$ . As the next claim shows, it also retains *some* of its soundness property, even with respect to assignments which are not necessarily feasible.

**Claim 3.3.** There exists a constant  $\alpha$ ,  $0 < \alpha < 1$ , such that  $\Psi_2$  has the following properties:

• Completeness: If  $\Psi$  is completely satisfiable, then  $\Psi_2$  is completely satisfiable as well.

Weakened Soundness: If Ψ is at most 2/|F|-satisfiable, then Ψ<sub>2</sub> is at most |F|<sup>-α</sup>-satisfiable (by any assignment).

To prove the claim we need the following proposition, showing that there cannot be many permissible LDFs for a domain – this implies that most local-readers in an LDF-reader will either reject or return the evaluation of one of a *short list* of permissible LDFs. This proposition appears in Section 4 as Claim 4.9, and is proven there.

**Proposition 3.4.** Let F be a domain, and let  $\rho > \left(\frac{r(F)}{|\mathcal{F}|}\right)^{c_g} d(F)$  where  $c_g$  is the same constant as in Lemma 3.2. Then for any assignment to F there can be at most  $2\rho^{-1}$   $\rho$ -permissible LDFs in all.

Proof of Claim 3.3:

**Completeness.** If  $\Psi$  is satisfiable, then there is a good assignment satisfying  $\Psi_{sc}$ . For each of the constructed LDF-readers, extend the assignment to its representation using the encoding-assignment of the associated domain. The extended assignment satisfies  $\Psi_2$ : A conjunction in  $\Psi_2$  contains localtests, which are all satisfied by encoding-assignments, and an equation  $\psi'$ .  $\psi'$ was generated from an equation  $\psi \in \Psi_{sc}$  by replacing variables with evaluators. But for encoding-assignments, the evaluators and the replaced variables have the same values. Hence since  $\psi$  is satisfied,  $\psi'$  is satisfied as well.

Weakened soundness. Fix an assignment  $\mathcal{A}$  for  $\Psi_2$ , and let  $\gamma$  be the fraction of conjunctions it satisfies. For an appropriate  $\alpha$ , we will show that if  $\gamma > |\mathcal{F}|^{-\alpha}$  then there exists a feasible assignment for  $\Psi_{sc}$  satisfying more than a  $\frac{2}{|\mathcal{F}|}$  fraction of its equation. This implies that  $\Psi$  is more than  $\frac{2}{|\mathcal{F}|}$ -satisfiable – a contradiction.

We denote  $a \doteq (1 - c_2)c_g/k$ , where  $c_2$  is the global constant mentioned in the Sum-Check Lemma (Lemma 3.1). Letting  $\rho \doteq |\mathcal{F}|^{-a}$ , it follows by the choice of a that  $\rho > (r/|\mathcal{F}|)^{c_g}d$ . Therefore by Lemma 3.2 we have that the LDF-readers have parameters  $(\rho, \epsilon)$ , where  $\epsilon = O(\rho^c)$  and c is as mentioned in the lemma.

An equation  $\psi \in \Psi_{sc}$  such that the fraction of satisfied conjunctions in  $\mathcal{E}_{\psi}$  is higher than  $k\epsilon$ , is said to be *potentially satisfiable*. Since the sets  $\mathcal{E}_{\psi}$  are all of the same size, it follows that the fraction of potentially satisfiable equations is at least  $\gamma - k\epsilon$ .

Consider a potentially satisfiable equation  $\psi$ .  $\mathcal{E}_{\psi}$  was generated from  $\psi$ by plugging in k LDF-readers  $\mathcal{R}_1, \ldots, \mathcal{R}_k$ , evaluating tuples in k domains  $F_1, \ldots, F_k$  respectively. A conjunction in  $\mathcal{E}_{\psi}$  is defined by choosing a localreader  $L_j$  out of each LDF-reader  $\mathcal{R}_j$ . For every j, the fraction of conjunctions in  $\mathcal{E}_{\psi}$  where  $L_j$  is  $\rho$ -erroneous is bounded by  $\epsilon$ , as implied by the parameters of the LDF-readers, and hence the fraction of conjunctions where *any* of the readers are erroneous is bounded by  $k\epsilon$ .

It follows that there exists a satisfied conjunction in  $\mathcal{E}_{\psi}$  in which no localreader is erroneous, namely the evaluator of each local-reader  $L_j$  gives the evaluations of a  $\rho$ -permissible LDF  $f_j$  with respect to the assignment of  $F_j$ . Hence if each domain  $F_j$  were re-assigned the function  $f_j$ ,  $\psi$  would be satisfied.

So far we have shown that the potentially satisfiable equations, which make at least a  $\gamma - k\epsilon$  fraction of the equations  $\psi \in \Psi_{sc}$ , can be satisfied by re-assigning the domains with  $\rho$ -permissible LDFs. For each domain F in  $\Psi_{sc}$ , choose a random  $\rho$ -permissible LDF, or the zero LDF if no such LDF exists, and re-assign it to F. We have obtained a *feasible assignment* for  $\Psi_{sc}$ . We compute the chance of a potentially satisfiable equation to be satisfied by the new assignment.

There are at most  $O(\rho^{-1}) \rho$ -permissible LDFs for each domain by Proposition 3.4, and each equation has variables from k domains. Hence the probability of a potentially satisfiable equation in  $\Psi_{sc}$  to be actually satisfied by the re-assignment is at least  $\Omega(\rho^k)$ . It follows that the expected fraction of satisfied equations in  $\Psi_{sc}$  is  $\Omega(\rho^k(\gamma - k\epsilon))$ , and hence at least one of the re-assignments achieves this fraction of satisfaction. We have thus shown that there exists a feasible assignment for  $\Psi_{sc}$  satisfying an  $\Omega(\rho^k(\gamma - k\epsilon))$  fraction of its equations.

We now choose a constant  $\alpha$  so that  $0 < \alpha < \min\{1 - ka, ac\}$  (note that such an  $\alpha$  exists). If  $\gamma > |\mathcal{F}|^{-\alpha}$ , then

$$\rho^{k}(\gamma - k\epsilon) = |\mathcal{F}|^{-ak}(\gamma - O(\mathcal{F}^{-ac})) \gg \frac{2}{|\mathcal{F}|}$$

hence there exists a feasible assignment for  $\Psi_{sc}$  satisfying more than a  $\frac{2}{|\mathcal{F}|}$  fraction of its equations.

### 3.3 Gap Amplification

The reduction now amplifies the soundness of  $\Psi_2$  by joining conjunctions together into larger conjunctions, generating  $\Psi_3$ . The soundness of  $\Psi_3$  is even stronger than needed, but it still has conjunctions rather than equations. The next subsection describes how conjunctions may be replaced by equations with only a small cost in the soundness, thus completing the reduction.

**The system**  $\Psi_3$ . Denote  $N \doteq \lceil 1/\alpha \rceil$ , where  $\alpha$  is the constant mentioned in Claim 3.3. The reduction generates  $\Psi_3$  by taking the conjunction of every ordered N-tuple of (not necessarily distinct) conjunctions from  $\Psi_2$ , that is

$$\Psi_3 = \{ \bigwedge_{i=1}^N \chi_i : \forall i \ \chi_i \in \Psi_2 \}$$

Note that it takes polynomial time in  $|\mathcal{F}|^N$ , and hence in n, to generate  $\Psi_3$ . Since each conjunction in  $\Psi_3$  is composed of a constant number of conjunctions from  $\Psi_2$ , the number of variables as well as the number of equations in each such conjunction is bounded by a constant. The next claim states the completeness and soundness properties of  $\Psi_3$ .

**Claim 3.5.**  $\Psi_3$  has the following properties:

- Completeness: If  $\Psi$  is completely satisfiable, then  $\Psi_3$  is completely satisfiable as well.
- Soundness: If  $\Psi$  is at most  $2/|\mathcal{F}|$ -satisfiable, then  $\Psi_3$  is at most  $1/|\mathcal{F}|$ -satisfiable.

The claim follows easily from Claim 3.3 and from the construction of  $\Psi_3$ .

## **3.4** From Conjunctions to Equations

 $\Psi_3$  is a system of conjunctions where, as mentioned above, the number of equations in each conjunction is bounded by a constant. We would like the reduction to transform it from a system of conjunctions into the final system  $\Psi'$  of quadratic-equations, but first we make sure that the number of equations in all the conjunctions of  $\Psi_3$  is the same. To do so the reduction adds equations of the form 0 = 0 where necessary.

The system  $\Psi'$ . To transform  $\Psi_3$  into  $\Psi'$ , the reduction replaces each conjunction in  $\Psi_3$  with the set of all linear-combinations over its equations (equations can be added or multiplied by a scalar, so the notion of a linear-combination of equations is well defined). Since the number of equations in each conjunction is constant the blow-up is polynomial in  $|\mathcal{F}|$ , and hence in n.

Since the number of variables in each conjunction of  $\Psi_3$  is bounded by a constant, the number of variables in each equation of  $\Psi'$  is constant as well. In order to complete the proof of Theorem 1, it is left to show that  $\Psi'$  has the soundness and completeness properties. This follows immediately from Claim 3.5 together with the next proposition.

**Proposition 3.6** (conjunction representation). Let  $\Psi_a$  be a system of conjunctions of equations over  $\mathcal{F}$ , where the number of equations in each conjunction is the same. Let  $\Psi_b$  be the system obtained from  $\Psi_a$  by replacing every conjunction  $\chi \in \Psi_a$  by all linear combinations over  $\mathcal{F}$  of its equations (with multiplicities, if the same equation occurs more than once). Then

- If  $\Psi_a$  is completely satisfied by a certain assignment, then the same assignment will satisfy  $\Psi_b$  as well.
- If  $\Psi_a$  is at most  $\gamma$ -satisfiable then  $\Psi_b$  is at most  $(\gamma + 1/|\mathcal{F}|)$ -satisfiable.

*Proof.* The first property is obvious from the definition of  $\Psi_b$ . To prove the second property, fix an assignment for the variables of  $\Psi_a$  and  $\Psi_b$ . Then it satisfies at most a  $\gamma$  fraction of the conjunctions in  $\Psi_a$ . For each conjunction  $\chi$  in  $\Psi_a$  denote by  $\omega(\chi)$  the fraction of equations replacing  $\chi$  that are satisfied in  $\Psi_b$ . Since each conjunction of  $\Psi_a$  is replaced by the same number of equations, the fraction of satisfied equations in  $\Psi_b$  is the average of  $\omega(\chi)$  over all the conjunctions  $\chi \in \Psi_a$ .

For a satisfied conjunction  $\chi$ ,  $\omega(\chi) = 1$ , and it is easy to observe that  $\omega(\chi) = 1/|\mathcal{F}|$  for any unsatisfied conjunction  $\chi$ . Since satisfied conjunctions make at most a  $\gamma$  fraction of the conjunctions in  $\Psi_a$ , we conclude that the fraction of satisfied equations in  $\Psi_b$  is no more than  $\gamma + 1/|\mathcal{F}|$ , as required.

# 4 The Composition-Recursion Constructor

This section contains the proof of the Composition-Recursion Constructor Lemma (Lemma 3.2), showing the constructor for Composition-Recursion LDF-readers, CR's for short. As a first step, we show a constructor for *restricted* LDF-readers, where some of the domains in the representation are considered *active*. These LDF-readers have good parameters only in the case where active domains are given feasible assignments. By a composition of several such LDF-readers we then get a CR.

**Definition 8** (restricted LDF-readers.). A restricted LDF-reader  $\mathcal{R}$  is an LDF-reader where some of the domains in the representation are considered active. The dimension, and the upper and lower degree parameters of all active domains must be the same. These parameters are called the active dimension, active upper-degree and active lower-degree of  $\mathcal{R}$ , and are denoted by  $d_*(\mathcal{R})$ ,  $r_*(\mathcal{R})$ , and  $s_*(\mathcal{R})$  respectively.

A local-reader L in  $\mathcal{R}$  may have variables from at most one active domain, which is called the active domain of L and is denoted by  $\text{Dom}_{\star}(L)$ .

**Parameters of restricted LDF-readers.** We measure the parameters of restricted LDF-readers only with respect to feasible assignments: An assignment for the representation of a restricted LDF-reader  $\mathcal{R}$  is said to be *active-feasible* if the assignment of every active domain is feasible (unlike in the case of equation-systems, we do not require the assignment of all domains to be feasible).  $\mathcal{R}$  is hence said to be a restricted ( $\rho, \epsilon$ )-LDF-Reader if for any active-feasible assignment, the fraction of  $\rho$ -erroneous local-readers is at most  $\epsilon$ . Note that in an encoding-assignment, all domains must still be given a good assignment.

Outline of this section. The next subsection shows a constructor for restricted LDF-readers (the definition of a constructor generalizes naturally for restricted LDF-readers), called Subspace-vs.-Point LDF-readers, SP's for short. These restricted LDF-readers are based upon the Plane-vs-Plane LDFreaders of [RS97]. The representation of an SP evaluating a tuple in a domain F contains, apart from F itself, only active domains, which have the same degree-parameters as F but a *constant* dimension parameter. Therefore, informally speaking, an SP LDF-reader uses constant-dimensional LDFs to represent an LDF over a space of higher dimension, and using evaluations of these constant-dimensional LDFs it produces consistent evaluations of the original LDF.

In Subsection 4.4 it is shown how the constant-dimensional active domains of an SP,  $\mathcal{R}$ , can be replaced by active domains that have non-constant dimension, but greatly decreased degree parameters. This allows the composition of other SP's over  $\mathcal{R}$ , as described in Subsection 4.5, to evaluate tuples in the replaced active domains. Subsection 4.6 shows how an iterative application of this procedure yields the Composition-Recursion LDF-reader (which is not restricted) and Subsection 4.7 proves its properties, thus completing the proof of Lemma 3.2.

### 4.1 Subspace-vs.-Point LDF-Readers

In this subsection we show the SP constructor – a constructor that generates Subspace-vs.-Point restricted LDF-readers. The representation of an SP that evaluates a k-tuple in a domain F contains, in addition to F, active domains with the same degree parameters as F but of dimension k + 2. Each domain is associated with a (k+2)-dimensional subspace U in  $\mathcal{F}^{d(F)}$ ; in an encodingassignment each of them is assigned the restriction to U of the LDF assigned to F. Before we go into the description of the constructor, let us state its properties in the following lemma.

**Lemma 4.1** (Subspace-vs.-Point LDF-reader). There exists a constructor that given a domain F and a k-tuple of points in  $\mathcal{F}^{d(F)}$ , generates a restricted LDF-reader  $\mathcal{R}$  as follows. The active domains of  $\mathcal{R}$  have the same upper and lower-degree parameters as F, and their dimension parameter equals k+2. Moreover,  $\mathcal{R}$  will have parameters  $(\rho, O(\rho^{1/3}))$  for all  $\rho$ 's which satisfy  $\rho > (r(F)/|\mathcal{F}|)^{c_g} d(F)$ , where  $0 < c_g \leq 1/2$  is a global constant<sup>‡</sup>.

### The Subspace-vs.-Point constructor

We now describe how the SP constructor generates an LDF-reader  $\mathcal{R}$ , given a domain F and a k-tuple  $(x_1, \ldots, x_k)$  of points in  $\mathcal{F}^{d(F)}$ . The SP constructor is used later as a procedure of the CR constructor, however in proving the parameters of the CR constructor we only rely on the properties that are stated in Lemma 4.1. Without loss of generality, throughout the construction

<sup>&</sup>lt;sup>‡</sup>This is the same constant as in Lemma 3.2, and in all other places where  $c_g$  appears

it is assumed that  $d(F) \ge k + 2$  – it is easy to adapt the construction for the case d(F) < k + 2.

The representation. Other than F itself, the representation only includes active domains, with upper-degree r(F), lower-degree s(F), and dimension k+2. The constructor first picks any (k-1)-dimensional affine subspace  $U_0 \subseteq \mathcal{F}^{d(F)}$  which contains all the points  $x_i$  of the tuple (if the  $x_i$ 's are in general position, there exists exactly one such subspace). Denote by  $\mathcal{S}ubSp(\mathcal{R})$  the set of (k + 2)-dimensional affine subspaces  $U \subseteq \mathcal{F}^{d(F)}$  which contain  $U_0$ . One active domain  $\mathcal{D}_U$  is then constructed for every affine subspace  $U \in \mathcal{S}ubSp(\mathcal{R})$ .

**Identification functions.** A good assignment  $\mathcal{A}$  to F assigns to it an [s(F), d(F)]-LDF f. In the encoding-assignment, the assignment to each domain  $\mathcal{D}_U$  represents the restriction of f to U. In order to represent  $f|_U$  as an LDF over  $\mathcal{F}^{k+2}$  the constructor chooses for each domain  $\mathcal{D}_U$  an arbitrary linear isomorphism  $\phi_U : U \to \mathcal{F}^{k+2}$ , called the *identification function* of U, that identifies each point  $y \in U$  with the point  $\phi_U(y)$  in  $\mathcal{F}^{k+2}$ .

**Encoding-assignments.** Let  $\mathcal{A}$  be a good assignment for F, assigning to it a [s(F), d(F)]-degree LDF f. The encoding-assignment for  $\mathcal{A}$  extends it by assigning to each domain  $\mathcal{D}_U$  the LDF  $f \circ (\phi_U^{-1})$ . Composing the assignment of  $\mathcal{D}_U$  with  $\phi_U$  therefore gives  $f|_U$ . Since  $\phi_U$  is linear,  $\mathcal{D}_U$  is assigned an s(F)-degree LDF, and hence the encoding-assignment of  $\mathcal{A}$  is a good assignment.

**Local-readers.** The SP constructor generates one local-reader for each domain  $\mathcal{D}_U$  and point  $y \in U$ . Its active domain is  $\mathcal{D}_U$ , its local-test is the single linear equation  $\mathcal{D}_U[\phi_U(y)] = F[y]$ , and for every  $1 \leq i \leq k$  its *i*'th evaluator is the term  $\mathcal{D}_U[\phi_U(x_i)]$ .

To get a better understanding of the structure of local-readers, fix an activefeasible assignment  $\mathcal{A}$  for the representation variables, and consider a localreader associated with a domain  $\mathcal{D}_U$  and a point y. The LDF assigned to  $\mathcal{D}_U$ represents an r(F)-degree LDF  $g_U$  over U, defined by  $g_U(x) \doteq \mathcal{A}(\mathcal{D}_U[\phi_U(x)])$ (this is the composition of the LDF assigned to  $\mathcal{D}_U$  with  $\phi_U$ ). The local-test therefore compares  $g_U(y)$  with the assignment of F[y], and the *i*'th evaluator returns  $g_U(x_i)$ . If  $\mathcal{A}$  is the encoding-assignment of an LDF f, then F[y] is assigned f(y), and  $g_U$  is the restriction of f to U. The local-test is hence satisfied in that case, and the values  $g_U(x_i)$  returned by the evaluators are in fact the values of f at the points  $x_i$ .

#### The SP constructor works.

It is easy to verify that the SP constructor indeed falls under the definition of a constructor. To verify the parameters of SP LDF-readers (which would conclude the proof of Lemma 4.1) consider an SP LDF-reader  $\mathcal{R}$ , that evaluates a k-tuple  $(x_1, \ldots, x_k)$  in a domain F, and fix an active-feasible assignment  $\mathcal{A}$  for its representation. As explained above,  $\mathcal{A}$  determines an r(F)-degree LDF  $g_U$  over every affine subspace  $U \in SubSp(\mathcal{R})$ .

The local-test of the local-reader determined by an affine subspace  $U \in SubSp(\mathcal{R})$  and a point  $y \in U$  verifies that  $g_U(y) = \mathcal{A}(F[y])$ , and its evaluators return the values of  $g_U$  at the points  $x_1, \ldots, x_k$ . The local-reader is  $\rho$ -erroneous if the local-test is satisfied, yet the values  $g_U(x_1), \ldots, g_U(x_k)$  are not the evaluation of any  $\rho$ -permissible LDF (with respect to the assignment of F) at  $x_1, \ldots, x_k$ . The next lemma bounds the fraction of erroneous localreaders, thus proving that  $\mathcal{R}$  has the parameters required in Lemma 4.1.

**Lemma 4.2** (SP parameters). For some global constant  $0 < c_g \leq 1/2$ , and for all  $\rho$ 's which satisfy  $\rho > (r(F)/|\mathcal{F}|)^{c_g} d(F)$ , the following holds.

Let U be a random affine subspace in  $SubSp(\mathcal{R})$ , and y be a random point in U (this determines a random local-reader). Let **E**rr be the event that  $g_U(y) = \mathcal{A}(F[y])$ , yet  $g_U(x_1), \ldots, g_U(x_k)$  are not the evaluation of any  $\rho$ -permissible LDF (with respect to the assignment of F) at  $x_1, \ldots, x_k$ . Then  $\Pr[\mathbf{E}rr] = O(\rho^{1/3})$ .

In fact we show an stronger statement than the above lemma. We bound by  $O(\rho^{1/3})$  the probability that  $g_U$  agrees with the assignment of F at y, yet  $g_U$  is not the restriction to U of any  $\rho$ -permissible LDF (it is easy to observe that this implies Lemma 4.2).

### 4.2 Subspace-vs.-Point Parameters

The proof of Lemma 4.2 is based on a lemma from [RS97] where a slightly different setting is discussed. While we are interested in the case where an LDF is associated with every (k + 2)-dimensional subspace in a certain

set  $SubSp(\mathcal{R})$ , the [RS97]-Lemma deals with the case where each plane (2dimensional affine subspace) is associated with an LDF defined over it.

**Definition 9** (plane-assignment). Suppose that every plane P in  $\mathcal{F}^{d(F)}$  is associated with an r(F)-degree LDF  $g_P$  over P. The correspondence  $P \to g_P$  is called a plane-assignment. An LDF  $g_P$  is called the plane-LDF assigned to P.

Another difference is that the [RS97]-Lemma discusses a different kind of permissibility, measured with respect to the plane-assignment instead of with respect to the assignment of F.

**Definition 10** (planewise-permissibility). Let  $(P \to g_P)$  be a plane-assignment. An r(F)-degree LDF f over  $\mathcal{F}^{d(F)}$  is said to be  $\rho$ -planewise-permissible if for at least a  $\rho$ -fraction of the planes P,  $g_P = f|_P$ .

A plane-LDF  $g_P$  is said to be  $\rho$ -planewise-permissible if it is the restriction to P of a  $\rho$ -planewise-permissible LDF f over  $\mathcal{F}^{d(F)}$ .

We now state the discussed lemma from [RS97]. It shows that planewisepermissibility can be tested by comparing the plane-LDFs assigned to two line-intersecting planes. If the plane-LDFs agree on the line then with high probability they are both planewise-permissible.

**Lemma 4.3** ([RS97]). There exists a global constant  $0 < c_g \leq 1/2$ , such that for any  $\rho > (r(F)/|\mathcal{F}|)^{c_g} d(F)$  the following holds.

Fix a plane-assignment  $(P \to g_P)$ . Let  $\ell$  be a random line in  $\mathcal{F}^{d(F)}$  and let  $P_1$  and  $P_2$  be two random, independently chosen planes that contain  $\ell$ . Denote by **E**rr the event that the plane-LDF assigned to  $P_1$  agrees on  $\ell$  with the plane-LDF assigned to  $P_2$ , yet they are not both  $\rho$ -planewise-permissible. Then  $\Pr[\mathbf{E}rr] = O(\rho)$ .

Starting from Lemma 4.3, we gradually approach Lemma 4.2 by a sequence of technical claims: Claim 4.4 shows Lemma 4.3 to hold even if two plane-LDFs are compared on a point rather than a line. Claim 4.5 deals with the case where a plane-LDF is compared against the assignment of F at a certain point, rather than against another plane-LDF. Claim 4.7 goes from planewise-permissibility to permissibility, showing that a random plane-LDF  $g_P$  that agrees with the assignment of F at a random point on P is with high probability the restriction to P of a  $\rho$ -permissible LDF. Finally Claim 4.11 completes the proof of Lemma 4.2 by showing that the same holds for LDFs  $g_{_U}$  associated with a random subspace  $U \in SubSp(\mathcal{R})$ , instead of plane-LDFs.

**Claim 4.4.** Fix a plane assignment  $(P \to g_P)$ , and suppose  $\rho$  satisfies the requirements of Lemma 4.3. Let y be a random point in  $\mathcal{F}^{d(F)}$ , let  $\ell$  be a random line containing it, and let  $P_1$  and  $P_2$  be random independently chosen planes that contain  $\ell$ . Denote by **E**rr the event that the plane-LDF assigned to  $P_1$  agrees on y with the plane-LDF assigned to  $P_2$ , yet they are not both  $\rho$ -planewise-permissible. Then  $\Pr[\mathbf{E}rr] = O(\rho)$ .

*Proof.* First note that y can be considered to be a random point on a randomly chosen line  $\ell$  in  $\mathcal{F}^{d(F)}$ , instead of the other way around.

The only case where  $\mathbf{E}rr$  occurs yet the event from Lemma 4.3 does not, is when the restrictions to  $\ell$  of the plane-LDFs of  $P_1$  and  $P_2$  differ, yet they agree on y. Since the restrictions to  $\ell$  are r(F)-degree LDFs, if they differ then the probability of agreement on the random point y is at most  $r(F)/|\mathcal{F}| \leq \rho$ . Hence by Lemma 4.3,  $\Pr[\mathbf{E}rr] \leq \rho + O(\rho) = O(\rho)$ .

The next step is to compare a plane-LDF to the assignment of F[y] for some point y on it, instead of to the value at y of another plane-LDF.

**Claim 4.5.** Fix a plane assignment  $(P \to g_P)$ , and suppose  $\rho$  satisfies the requirements of Lemma 4.3. Let P be a random plane and y be a random point on P. Denote by **E**rr the event that  $g_P(y) = \mathcal{A}(F[y])$ , yet  $g_P$  is not  $\rho$ -planewise-permissible. Then  $\Pr[\mathbf{E}rr] = O(\rho^{1/2})$ .

*Proof.* To be able to apply Claim 4.4, we redefine y and P, and introduce new random variables as follows. Let y be a random point in  $\mathcal{F}^{d(F)}$ ,  $\ell$  be a random line containing y, and P and P' be random independently chosen planes that contain  $\ell$  (note that to obtain the claim it is enough to prove a bound on  $\Pr[\mathbf{E}rr]$  in these settings). Let  $\mathbf{E}rr_2$  be the event that  $g_P$  and  $g_{P'}$ agree on y yet they are not both  $\rho$ -planewise-permissible. Claim 4.4 implies that the probability of  $\mathbf{E}rr_2$  is bounded by  $O(\rho)$ .

To use the bound we have for  $\mathbf{E}rr_2$  we first show that for every fixed line  $\ell_0$ and point  $y_0 \in \ell_0$ ,

$$\Pr[\mathbf{E}rr|\ell = \ell_0, y = y_0] \le \left(\Pr[\mathbf{E}rr_2|\ell = \ell_0, y = y_0]\right)^{1/2} \tag{1}$$

Let  $\mathbf{E}rr'$  be the event that  $g_{P'}(y) = \mathcal{A}(F[y])$  yet  $g_{P'}$  is not  $\rho$ -planewisepermissible (it is similar to  $\mathbf{E}rr$ , only for P' instead of P). Obviously

$$\Pr[\mathbf{E}rr_2|\ell=\ell_0, y=y_0] \ge \Pr[\mathbf{E}rr \wedge \mathbf{E}rr'|\ell=\ell_0, y=y_0]$$

because the event on the left-hand side contains the one on the right-hand side. Since P and P' are independently chosen given  $\ell_0$ , we have

$$\Pr[\mathbf{E}rr \wedge \mathbf{E}rr' | \ell = \ell_0, y = y_0] = \Pr[\mathbf{E}rr | \ell = \ell_0, y = y_0] \cdot \Pr[\mathbf{E}rr' | \ell = \ell_0, y = y_0]$$
$$= \left(\Pr[\mathbf{E}rr | \ell = \ell_0, y = y_0]\right)^2$$

which together with the above inequality implies Equation 1.

One may discard the conditioning in Equation 1, obtaining

$$\Pr[\mathbf{E}rr] \le \Pr[\mathbf{E}rr_2]^{1/2}$$

using the law of complete probability and the concavity of the square-root function. Since the probability of  $\mathbf{E}rr_2$  is bounded by  $O(\rho)$ , this obtains the claim.

Our next step is to convert the statement of Claim 4.5 from planewisepermissibility to permissibility in the usual sense. This requires the following bound on the number of planewise-permissible LDFs.

**Claim 4.6.** Fix a plane assignment  $(P \rightarrow g_P)$ , and suppose  $\rho$  satisfies the requirements of Lemma 4.3. Then the number of  $\rho$ -planewise-permissible LDFs is less than  $2\rho^{-1}$ .

*Proof.* The proof is similar to that of Claim 4.9 below.

We can now prove the analogue of Claim 4.5 for permissibility in the usual sense.

**Claim 4.7.** Fix a plane assignment  $(P \to g_P)$ , and suppose  $\rho$  satisfies the requirements of Lemma 4.3. Let P be a random plane and y be a random point on P. Denote by Err the event that  $g_P(y) = \mathcal{A}(F[y])$ , yet there is no  $\rho$ -permissible LDF f (with respect to the assignment of F), such that  $g_P = f|_P$ . Then  $\Pr[\mathbf{E}rr] = O(\rho^{1/3})$ .

*Proof.* We separate  $\mathbf{E}rr$  into two events, and bound the probability of each by  $O(\rho^{1/3})$ : Let  $\mathbf{E}rr_1$  be the event where  $\mathbf{E}rr$  occurs and in addition  $g_P$  is **not**  $\rho^{2/3}$ -planewise-permissible; and let  $\mathbf{E}rr_2$  be the event where  $\mathbf{E}rr$  occurs and in addition  $g_P$  is  $\rho^{2/3}$ -planewise-permissible.

By applying Claim 4.5 using  $\rho^{2/3}$  instead of  $\rho$ , we obtain that the probability of  $\mathbf{E}rr_1$  is bounded by  $O(\rho^{1/3})$  as required (since  $\rho > (r(F)/|\mathcal{F}|)^{c_g} d(F)$ as required in Lemma 4.3,  $\rho^{2/3}$  satisfies this requirement as well).

It is left to bound the probability of  $\mathbf{E}rr_2$ . By definition it occurs only when  $g_P(y) = \mathcal{A}(F[y])$  and there exists a  $\rho^{2/3}$ -planewise-permissible LDF fwhich is not  $\rho$ -permissible, such that  $g_P = f|_P$ . For an LDF f over  $\mathcal{F}^{d(F)}$ , denote by  $\mathbf{E}rr_3(f)$  the event where  $g_P(y) = \mathcal{A}(F[y])$  and  $g_P = f|_P$  (note that this implies  $f(y) = \mathcal{A}(F[y])$ ). Then the probability of  $\mathbf{E}rr_2$  is bounded by the sum of  $\Pr[\mathbf{E}rr_3(f)]$  over all  $\rho^{2/3}$ -planewise-permissible LDFs f which are not  $\rho$ -permissible.

Let us bound the probability of  $\mathbf{E}rr_3(f)$  for such an LDF f. Since f is not  $\rho$ -permissible the probability that it satisfies  $f(y) = \mathcal{A}[F(y)]$  is bounded by  $\rho$ , because y is a uniformly random point in  $\mathcal{F}^{d(F)}$ . The probability of  $\mathbf{E}rr_3(f)$  is therefore bounded by  $\rho$  as well. Since f should be  $\rho^{2/3}$ -planewise-permissible, there can be at most  $2\rho^{-2/3}$  such f's by Claim 4.6, and therefore a bound of  $2\rho^{-2/3}\rho = O(\rho^{1/3})$  is obtained for the probability of  $\mathbf{E}rr_2$ .

Note that the statement of Claim 4.7 is similar to what we wish to establish (see the remark following Lemma 4.2), only for planes rather than (k+2)-dimensional subspaces in  $SubSp(\mathcal{R})$ . Claim 4.11 proves that by considering a random plane  $P, y \in P \subseteq U$ , in addition to the random subspace U and the random point  $y \in U$ . Claim 4.7 can be applied to P to obtain Claim 4.11, but for it to be applicable it should be shown that when a random subspace  $U \in SubSp(\mathcal{R})$  is chosen , and then a random plane P contained in U, and then a point  $y \in P, P$  and y are almost uniformly distributed. This is shown in the following claim.

**Claim 4.8.** Let U be a random subspace in  $SubSp(\mathcal{R})$ , and let P be a random plane contained in U and y a random point in P. The distribution of P and y is almost uniform, that is if Plns denotes the set of planes in  $\mathcal{F}^{d(F)}$  then

$$\sum_{P_0 \in \mathcal{P} \ln s} |\Pr(P = P_0) - |\mathcal{P} \ln s|^{-1}| \le O(|\mathcal{F}|^{-1})$$

and

$$\sum_{y_0 \in \mathcal{F}^{d(F)}} |\Pr(y = y_0) - |\mathcal{F}|^{-d(F)}| \le O(|\mathcal{F}|^{-1})$$

*Proof.* We begin by proving the second inequality. Observe that since all the subspaces in  $SubSp(\mathcal{R})$  contain  $U_0$  the probability of the random point y in U to yield a specific point in  $U_0$  is higher than  $|\mathcal{F}|^{-d(F)}$ , the probability of a uniformly random point. Also, the probability of y to yield a point outside of  $U_0$  is smaller than  $|\mathcal{F}|^{-d(F)}$ . Hence

$$\begin{split} &\sum_{y_0 \in \mathcal{F}^{d(F)}} |\Pr[y = y_0] - |\mathcal{F}|^{-d(F)}| \\ &= \sum_{y_0 \in U_0} (\Pr[y = y_0] - |\mathcal{F}|^{-d(F)}) + \sum_{y_0 \in \mathcal{F}^{d(F)} \setminus U_0} (|\mathcal{F}|^{-d(F)} - \Pr[y = y_0]) \\ &= 2 \sum_{y_0 \in U_0} (\Pr[y = y_0] - |\mathcal{F}|^{-d(F)}) \qquad (\text{since probabilities sum up to } 1) \\ &< 2 \Pr[y \in U_0] = 2|U_0|/|U| = 2|\mathcal{F}|^{-3} = O(|\mathcal{F}|^{-1}) \,, \end{split}$$

thus obtaining the second inequality. The proof of the first inequality uses similar arguments, however it is more tedious. We therefore only sketch it here.

Two main observations are needed to prove the first inequality. First, one needs to observe that if three random points  $y_1, y_2, y_3$  are independently chosen within U, then their joint distribution is within statistical distance  $O(|\mathcal{F}|^{-1})$  from the distribution of three points chosen independently from  $\mathcal{F}^{d(F)}$ . This follows by applying the argument used to prove the second inequality three consecutive times. At each application, U is conditioned on containing the affine space generated by  $U_0$  and the previously selected points.

The second observation, which is an easy exercise, is that the distribution of the affine span of three independently chosen points in  $\mathcal{F}^{d(F)}$ , is within statistical distance  $O(|\mathcal{F}|^{-1})$  from the distribution of a random plane in  $\mathcal{P}lns$ . Together with the first observation we obtain the desired inequality.

Before we move to the final claim, we need the following two bounds. Claim 4.9, which appears in Section 3 as Proposition 3.4, bounds number of  $\rho$ -permissible LDFs. Claim 4.10 bounds the fraction of planes on which two distinct LDFs may agree. **Claim 4.9.** Suppose  $\rho$  satisfies the requirements of Lemma 4.3. Then there are less than  $2/\rho$   $\rho$ -permissible LDFs.

Proof. Assume for the sake of contradiction that there exists a set  $\mathcal{P}er$  containing  $2/\rho$  distinct  $\rho$ -permissible LDFs. For each LDF  $f \in \mathcal{P}er$  denote by U(f) the set of points  $y \in \mathcal{F}^{d(F)}$  such that f is the only LDF in  $\mathcal{P}er$  satisfying  $f(y) = \mathcal{A}(F[y])$ .

Each LDF  $f \in \mathcal{P}er$  is  $\rho$ -permissible, hence it agrees with  $\mathcal{A}(F)$  on at least a  $\rho$ -fraction of the points. We bound from above the fraction of points for which it also agrees with other LDFs in  $\mathcal{P}er$ . Any other LDF in  $\mathcal{P}er$  agrees with f on at most an  $\frac{r(F)}{|\mathcal{F}|}$  fraction of the points, so overall f may agree with other LDFs in  $\mathcal{P}er$  on at most a  $\frac{2r(F)}{\rho|\mathcal{F}|}$  fraction of the points. From the assumption  $\rho > \left(\frac{r(F)}{|\mathcal{F}|}\right)^{c_g} d(F)$  it follows in particular that  $\rho^2 > 4r(F)/|\mathcal{F}|$ (recall that  $c_g < 1/2$  and that we assume  $d(F) \ge k + 2 > 2$ ), and therefore  $\frac{2r(F)}{\rho|\mathcal{F}|} < \frac{1}{2}\rho$ .

f thus agrees with  $\mathcal{A}(F)$  on at least  $\rho$  of the points, and on less than a  $\frac{1}{2}\rho$  fraction of the points it agrees with other LDFs in  $\mathcal{P}er$ . It follows that for every  $f \in \mathcal{P}er$ , U(f) contains more than a  $\frac{1}{2}\rho$  fraction of the points. Since the sets U(f) are disjoint, it follows that the fraction of all points contained in any of the U(f)'s is greater than  $\frac{1}{2}\rho \cdot |\mathcal{P}er| \geq 1$ . This is a contradiction.

**Claim 4.10.** For any t > 0, two distinct [r, t]-LDFs must disagree on all but at most an  $r/|\mathcal{F}|$  fraction of their possible restrictions to planes.

*Proof.* Let f and g be two distinct r-degree LDFs over  $\mathcal{F}^t$ , and let P be a random plane in  $\mathcal{F}^t$ . We are to evaluate the probability that  $f|_P$  equals  $g|_P$ . Let y be a random point on P. y is uniformly distributed in  $\mathcal{F}^t$ , and therefore it produces a disagreement with probability at least  $1 - \frac{r}{|\mathcal{F}|}$  (this is a well known property of LDFs). Since y can produce a disagreement only in the case that there is a disagreement over P, it implies that there is a disagreement over P with probability at least  $1 - \frac{r}{|\mathcal{F}|}$ .

The following claim directly implies Lemma 4.2.

**Claim 4.11.** Suppose  $\rho$  satisfies the requirements of Lemma 4.3. Let U be a random affine subspace in  $SubSp(\mathcal{R})$ , and y be a random point in U. Let **E**rr be the event that  $g_U(y) = \mathcal{A}(F[y])$ , yet there is no  $\rho$ -permissible LDF (with respect to the assignment of F) whose restriction to U gives  $g_U$ . Then  $\Pr[\mathbf{E}rr] = O(\rho^{1/3})$ . *Proof.* Let P be a random plane contained in the random subspace U. Without loss of generality, we may assume that y is a random point in P.

We define two events  $\mathbf{E}rr_1$  and  $\mathbf{E}rr_2$  such that  $\mathbf{E}rr_1 \cup \mathbf{E}rr_2$  contains  $\mathbf{E}rr$ , and bound the probability of each by  $O(\rho^{1/3})$ . Let  $\mathbf{E}rr_1$  be the event that  $g_U(y) = \mathcal{A}(F[y])$ , yet there is no  $\rho$ -permissible LDF f that agrees with  $g_U$  on P, namely  $f|_P = g_U|_P$ . Let  $\mathbf{E}rr_2$  be the event that there is no  $\rho$ -permissible LDF whose restriction to U gives  $g_U$ , yet there exists such an LDF that agrees with  $g_U$  on P. Obviously  $\mathbf{E}rr \subseteq \mathbf{E}rr_1 \cup \mathbf{E}rr_2$ .

**Bounding**  $\Pr[\mathbf{E}rr_2]$ . For a  $\rho$ -permissible LDF f, let  $\mathbf{E}rr_3(f)$  be the event that  $f|_U \neq g_U$ , yet  $f|_P = g_U|_P$ .  $\mathbf{E}rr_2$  is contained in the union of the events  $\mathbf{E}rr_3(f)$  over all  $\rho$ -permissible LDFs f. For a  $\rho$ -permissible LDF f,

$$\Pr[\mathbf{E}rr_3(f)|U=V] \le r(F)/|\mathcal{F}|$$

for every fixed subspace  $V \in SubSp(\mathcal{R})$ , by applying Claim 4.10 to V. It follows that  $\Pr[\mathbf{E}rr_3(f)]$  is bounded by  $r(F)/|\mathcal{F}|$  as well. Since there are less than  $2/\rho \rho$ -permissible LDFs in all, we obtain that

$$\Pr[\mathbf{E}rr_2] < \frac{2r(F)}{\rho|\mathcal{F}|} < \rho$$

(the last inequality follows easily from the restriction on  $\rho$ ).

**Bounding**  $\Pr[\mathbf{E}rr_1]$ . We change the distribution of U, P and y, by letting P be a random plane in  $\mathcal{F}^{d(F)}$ , U be a random space in  $\mathcal{S}ubSp(\mathcal{R})$  that contains P, and y be a random point in P. By Claim 4.8, the statistical distance between the new distribution of P, which is uniform, and the original distribution is  $O(|\mathcal{F}|^{-1})$ . Under both the original and the new distributions, the distribution of U and y conditioned on P being a fixed plane  $P_0$  are the same -U is a random space in  $\mathcal{S}ubSp(\mathcal{R})$  that contains  $P_0$  and y is a random point in  $P_0$ . It follows that the statistical distance between the new joint distribution of U, P and y, and the original distribution is bounded by  $O(|\mathcal{F}|^{-1})$ . It is hence enough to bound the probability of  $\mathbf{E}rr_1$  according to the new distribution.

Let  $g_P \doteq g_U|_P$  be considered as a random plane-assignment for the (random) plane P. The definition of  $\mathbf{E}rr_1$  can hence be articulated as the event that  $g_P(y) = \mathcal{A}(F[y])$ , yet there is no  $\rho$ -permissible LDF f such that  $g_P = f|_P$ . Claim 4.7 naturally extends to the case where the planeassignments is random as long as the assignment to F is not random, hence it implies that  $\Pr[\mathbf{E}rr_2] = O(\rho^{1/3})$  (note that P is a uniformly random plane and y is a random point in P).

### 4.3 Overview of the CR-Constructor

Let us give an overview of the CR (Composition-Recursion) constructor. Given a domain and a tuple, the CR constructor generates a constant-length sequence of restricted LDF-readers that ends with the final, unrestricted, CR LDF-reader. Each transformation of an LDF-reader  $\mathcal{R}$  in the sequence into the next (except for the final one) has the same two steps as follows.

**Extension.** In the first step, an *extension-procedure* is applied to each active domain of  $\mathcal{R}$ , replacing it by a domain with greatly reduced degree parameters in the price of an increased dimension parameter. The active degree and dimension parameters of  $\mathcal{R}$  are thus changed, but its other properties are maintained.

**Composition.** The second step is the application of the *composition* procedure, which incorporates new LDF-readers into  $\mathcal{R}$ . First, it generates several new LDF-readers using the SP constructor (actually, any constructor with properties as in Lemma 4.1 will do), applying it to different active domains and tuples. The domains generated in the process then become active instead of the old active domains. These new active domains are of constant dimension, and because of the extension step their degree parameters are greatly reduced with respect to the active domains of  $\mathcal{R}$ . Finally the newly generated local-readers are plugged into the local-readers of  $\mathcal{R}$ , generating the next LDF-reader in the sequence.

We proceed as follows. First, in the next subsection, we give a formal definition of an extension and show the two extension-procedures used by the CR constructor. In Subsection 4.5 we describe the composition procedure and prove its properties. In Subsection 4.6 we describe the CR constructor and then we prove its correctness in Subsection 4.7.

### 4.4 Extensions

An extension of a domain F is a domain G which *contains* the variables of F: Each variable F[x] is endowed with another name  $G[\phi(x)]$ . The function  $\phi : \mathcal{F}^{d(F)} \to \mathcal{F}^{d(G)}$  is called the *gluing* of F to G. The extension must preserve good and feasible assignments as follows.

**Definition 11** (extension). Let F be a domain, and let G be a domain that contains the variables of F. G is called an extension of F if the following properties hold:

- Extension Property: Any good assignment  $\mathcal{A}$  for F can be extended to a good assignment for G, called the encoding-assignment or the encoding LDF of  $\mathcal{A}$ .
- Restriction Property: The restriction to F of any feasible assignment for G is a feasible assignment for F.

The point about extensions is that they allow the representation of an LDF assigned to a domain F by an encoding LDF with different properties. We can hence replace the active domains of a restricted  $(\rho, \epsilon)$ -LDF-reader  $\mathcal{R}$  by their extensions and obtain a restricted  $(\rho, \epsilon)$ -LDF-reader where the active degree parameters are different, usually considerably smaller.

**Proposition 4.12** (extension). Let  $\mathcal{R}$  be a restricted  $(\rho, \epsilon)$ -LDF-reader, evaluating a tuple in a domain F. Suppose that for each active domain G of  $\mathcal{R}$ , e(G) is an extension of G, and that all extensions have the same parameters. Then the LDF-reader  $\mathcal{R}'$  obtained from  $\mathcal{R}$  by just declaring these extensions as the active domains of  $\mathcal{R}'$  is a restricted  $(\rho, \epsilon)$ -LDF-reader.

*Proof.* It is given that all the active domains of  $\mathcal{R}'$  have the same parameters. To show that  $\mathcal{R}'$  is a valid restricted LDF-reader we define an encoding-assignment of  $\mathcal{R}'$ , for every good assignment to F.

Given a good assignment for F, let  $\mathcal{A}$  be its encoding-assignment with respect to  $\mathcal{R}$ . For each active domain G of  $\mathcal{R}$ , assign the encoding-assignment of  $\mathcal{A}(G)$  to its extension e(G). This obtains a good assignment for the representation of  $\mathcal{R}'$ , and since the assignments to the variables of  $\mathcal{R}$  are not changed all local-tests are satisfied and all local-readers return evaluations consistent with the assignment of F.

The fact that  $\mathcal{R}'$  has parameters  $(\rho, \epsilon)$  follows easily from the restriction property of each extension e(G), which implies that the restriction of an active-feasible assignment for  $\mathcal{R}'$  yields an active-feasible assignment for  $\mathcal{R}$ .

**Extension-procedures.** An extension-procedure is an algorithm which given a domain F, generates an extension G of F. The running time of the algorithm must be polynomial in  $|\mathcal{F}|^{d(G)}$ . We next show the two extension-procedures used by the CR constructor – the power-substitution and the linearization extension-procedures. In the sequence of restricted LDF-readers that is generated by the CR constructor (see the overview above), the power-substitution extension-procedure is used in the generation of all restricted LDF-readers but the last. The last restricted LDF-reader is generated using the linearization extension-procedure, and thus has active domains with lower-degree and upper-degree 1. The final, unrestricted, CR LDF-reader is obtained by replacing each such domain with variables that represent the coefficients of a linear function.

Given a domain F, the power-substitution extension-procedure constructs an extension G with greatly reduced degree parameters in the price of increasing the dimension parameter. The linearization extension-procedure, when applied to a domain F, yields a domain G with lower-degree and upper-degree 1. The dimension of G is, however, exponential in the degree parameters of F, hence the linearization is applied by the CR constructor only after very small active degree parameters are achieved.

**Gap consumption.** Recall that if G is an extension of a domain F, then for every good assignment to F there must be a good assignment for Gwhich extends it. This may (and in fact does, in all cases discussed herein) impose a lower-bound on the lower-degree of G. Similarly, the restriction property of extensions yields an upper-bound on the upper-degree of G. In the case of the power-substitution extension-procedure, this forces the gap between the upper and lower-degrees of G to be smaller than for F. That is, if G is the extension of a domain F obtained by the power-substitution extension-procedure, then r(G)/s(G) < r(F)/s(F).

Since the CR constructor applies the power-substitution extension-procedure several times as described in the overview, if it is applied to a domain Fwhere the gap between the upper-degree and lower-degree is not large enough (see Lemma 3.2), domains are eventually created where the upper-degree is smaller than the lower-degree. Since the linearization extension-procedure is not applicable to such domains, the CR constructor would not be able to construct an LDF-reader for F.

#### The power-substitution extension-procedure

We begin by stating the properties of the power-substitution extensionprocedure. For simplicity, we omit floor and ceiling signs where they are not essential.

#### **Proposition 4.13** (power-substitution). There exists an

extension-procedure, called power-substitution, which given a domain F and a parameter b > 1, generates an extension G of F with the following parameters: For  $t \doteq \lceil \log_b(s(F) + 1) \rceil$ ,

- d(G) = d(F)t
- s(G) = d(F)t(b-1)
- $r(G) = r(F)/b^{t-1} (\geq r(F)/s(F))$

The procedure is based on the idea that by replacing powers of variables in an LDF f with new auxiliary variables, the degree of f may be decreased dramatically. For example, we fix an LDF over one variable  $f(u_1) = u_1^{12} + u_1^{25}$ (the handling of multi-variate LDFs is very similar), and show an encoding LDF g over three variables.

g is obtained from f by substituting powers of  $u_1$  with new variables. Informally speaking, if  $v_0$  is considered as representing  $u_1$ ,  $v_1$  is considered as representing  $u_1^3$ , and  $v_2$  – as representing  $u_1^9$ , then  $u_1^{12} = v_1v_2$ , and  $u_1^{25} = v_0v_1^2v_2^2$  (note that we used the base 3 representation of 12 and 25). Replacing these terms in f obtains an LDF  $g(v_0, v_1, v_2) = v_1v_2 + v_0v_1^2v_2^2$  of degree 5 rather than 25. g encodes and extends f in the sense that  $g(u_1, u_1^3, u_1^9) = f(u_1)$  for every  $u_1 \in \mathcal{F}$ .

For a domain G, obtained from a domain F using the power-substitution extension-procedure with parameter b, an LDF f of degree s(F) assigned to F is encoded by an LDF g over  $\mathcal{F}^{d(G)}$  as follows. g is obtained from f by taking an auxiliary variable for each power of the form  $u_i^{b^e}$  of a variable  $u_i$ of f. Any other power  $u_i^j$  of  $u_i$  can then be replaced by a monomial over the new variables of degree at most b-1 in each variable, using the base-brepresentation of j. *Proof of Proposition 4.13:* We begin by describing the power-substitution extension-procedure and then prove that it has the required properties.

**The procedure.** Given a domain F and a parameter b, the procedure first generates a domain G with parameters as stated in the proposition. It then generates a gluing function  $\phi : \mathcal{F}^{d(F)} \to \mathcal{F}^{d(G)}$  as follows: For every  $x = (u_1, \ldots, u_{d(F)}) \in \mathcal{F}^{d(F)}, \phi(x)$  is defined to be

$$(u_1, u_1^b, u_1^{b^2}, \dots, u_1^{b^{t-1}}, u_2, u_2^b, u_2^{b^2}, \dots, u_2^{b^{t-1}}, \dots, u_{d(F)}, \dots, u_{d(F)}^{b^{t-1}}) \in \mathcal{F}^{d(G)}$$

Finally, each variable of the form  $G[\phi(x)]$  is discarded, and the name  $G[\phi(x)]$  is endowed to the variable F[x] (which now has more than one name).

It is clear that the above procedure generates a domain G with the required parameters, in time polynomial in  $|\mathcal{F}|^{d(G)}$ . It remains to show that G is indeed an extension of F, namely that it has the extension and restriction properties.

**Extension property.** Suppose F is assigned an [s(F), d(F)]-LDF f (namely a good assignment). We now show its encoding LDF g – it should be an [s(G), d(G)]-LDF satisfying  $g \circ \phi = f$ , so that when assigned to G it does not conflict with F. First, let  $f(u_1, \ldots, u_{d(F)})$  be written as a polynomial formula P over the variables  $u_1, \ldots, u_{d(F)}$ . P is transformed into a polynomial formula P' over the variables

$$v_{(1,0)}, v_{(1,1)}, \dots, v_{(1,t-1)}, v_{(2,0)}, v_{(2,1)}, \dots, v_{(2,t-1)}, \dots, v_{(d(F),0)}, \dots, v_{(d(F),t-1)}$$

by replacing each term  $u_i^j$  in P with a monomial  $m_{(i,j)}$  over  $v_{(i,0)}, ..., v_{(i,t-1)}$ : Since the term  $u_i^j$  appears in P we gather that  $j \leq s(F)$ , and hence its representation as a number in base b has at most t digits. Let  $e_{t-1}, \ldots, e_1, e_0$ be the base b representation of j, and let

$$m_{(i,j)} \doteq (v_{(i,0)})^{e_0} (v_{(i,1)})^{e_1} \dots (v_{(i,t-1)})^{e_{t-1}}$$

Replacing each term  $u_i^j$  in P with the monomial  $m_{(i,j)}$  we obtain P', and then we define g by

$$g(v_{(i,0)},\ldots,v_{(d(F),t-1)}) \doteq P'(v_{(i,0)},\ldots,v_{(d(F),t-1)})$$
Since each monomial  $m_{(i,j)}$  is of degree at most t(b-1), it easily follows that g is an [s(G), d(G)]-LDF. Considering  $m_{(i,j)}$  as a function over  $\mathcal{F}^{d(G)}$ , it is also easy to see that for all  $(u_1, \ldots, u_{d(F)})$ ,  $(m_{(i,j)} \circ \phi)(u_1, \ldots, u_{d(F)}) = u_i^j$ . It follows that  $g \circ \phi = f$ , and hence g is indeed an encoding-LDF.

**Restriction property.** Suppose G is given a feasible assignment, namely it is assigned an [r(G), d(G)]-LDF g. The restriction of the assignment to F is hence an LDF f over  $\mathcal{F}^{d(F)}$ , given by  $f = g \circ \phi$ . The degree of f is at most  $\deg(f) = \deg(g) \deg(\phi) = r(G)b^{t-1} = r(F)$ . The restriction is hence a feasible assignment for F, as required.

#### The linearization extension-procedure

The linearization extension-procedure is very similar to the power-substitution procedure. The idea is to encode an LDF f by a *linear* LDF, replacing every monomial by a new auxiliary variable (recall that in the power-substitution, auxiliary variables where only introduced for some powers of variables in f). Since many auxiliary variables are used, the dimension increases dramatically.

**Proposition 4.14** (linearization). There exists an extension-procedure called linearization, which given a domain F with  $s(F) \leq r(F)$ , generates an extension G with the following parameters: For  $t \doteq \binom{s(F)+d(F)}{d(F)}$ ,

- d(G) = t
- s(G) = 1
- r(G) = 1

*Proof.* We begin by describing the linearization extension-procedure, and then prove that it has the required properties.

**The procedure.** Given a domain F, the procedure first generates a domain G with parameters as stated above. To generate the gluing function, the procedure picks an arbitrary enumeration  $m_1, \ldots, m_t$  of the monomial functions of degree at most s(F) over  $\mathcal{F}^{d(F)}$  (note that there are exactly t such monomials). The gluing function  $\phi : \mathcal{F}^{d(F)} \to \mathcal{F}^{d(G)}$  is then defined by

$$\forall x \in \mathcal{F}^{d(F)} \qquad \phi(x) \doteq (m_1(x), \dots, m_t(x))$$

Having defined the gluing function, F and G are then "glued" in the usual way – each variable of the form  $G[\phi(x)]$  is discarded, and the name  $G[\phi(x)]$  is endowed to the variable F[x] (which now has more than one name).

The procedure clearly generates a domain G with the required parameters, in time polynomial in  $|\mathcal{F}|^{d(G)}$ . It remains to verify that G has the extension and restriction properties.

**Extension property.** Suppose F is assigned an [s(F), d(F)]-LDF f, and let us construct its encoding LDF – a linear LDF over  $\mathcal{F}^{d(G)}$  satisfying  $g \circ \phi = f$ . First, one can write f as a linear-combination of the monomial functions of degree at most s(F):

$$f = \sum_{i=1}^{t} \gamma_i m_i$$

g is then defined by

$$\forall (v_1, \dots, v_t) \in \mathcal{F}^{d(G)} \qquad g(v_1, \dots, v_t) \doteq \sum_{i=1}^t \gamma_i v_i$$

It is clear that  $g \circ \phi = f$ , as desired.

**Restriction property.** Suppose G is given a feasible assignment, namely it is assigned a linear LDF g. The restriction of the assignment to F is the LDF  $f = g \circ \phi$ . Since  $\phi$  is of degree s(F) and g is linear, the degree of f is at most  $s(F) \leq r(F)$ , as required.

# 4.5 The Composition Procedure

We now turn to describe the composition procedure of the CR constructor algorithm. It takes as input a restricted LDF-reader  $\mathcal{R}$ , and generates a restricted LDF-reader  $\mathcal{R}'$  with the same active degree parameters, but where the dimension of the active domains is constant.

Suppose an LDF-reader  $\mathcal{R}$  is given, which evaluates a tuple  $(u_1, \ldots, u_k)$  in a domain F. The composition procedure has two main steps: First it generates new LDF-readers using the SP constructor as a sub-procedure, and then it incorporates them into  $\mathcal{R}$ .

**Uniformization.** Recall that each local-reader L of  $\mathcal{R}$  has variables from only one active domain,  $\text{Dom}_{\star}(L)$ . Before applying the main two steps, it is convenient to make sure that all local-readers L in  $\mathcal{R}$  have the same number of *active variables*, namely variables from  $\text{Dom}_{\star}(L)$ . Denoting the maximal number of active variables in a local-reader of  $\mathcal{R}$  by t, the composition procedure adds arbitrary variables to local-readers so that all have t active variables (the variables may be added anywhere in the local-reader, with zero coefficients).

**Generating new LDF-readers.** For each local-reader L in  $\mathcal{R}$ , the procedure now generates an LDF-reader denoted  $\mathcal{R}_L$  as follows. If G is the active domain of L and  $G[x_1], \ldots, G[x_t]$  are its active variables, then  $\mathcal{R}_L$  is generated by calling the SP constructor (see Lemma 4.1) with parameters G and  $(x_1, \ldots, x_t)$ .

**Domain incorporation.** The composition procedure now incorporates the domains of the new LDF-readers into  $\mathcal{R}$ : The newly generated domains are added to the representation. The active domains of  $\mathcal{R}$  cease to be active – the active domains of  $\mathcal{R}'$  are the active domains of the newly generated LDF-readers.

**Local-reader incorporation.** In an active-feasible assignment for  $\mathcal{R}'$ , the active variables of  $\mathcal{R}$ -local-readers L are no longer promised to be assigned the evaluation of a single feasible LDF over  $\text{Dom}_{\star}(L)$ . These variables are therefore replaced by the evaluators of local-readers of  $\mathcal{R}_L$ , which supposedly return evaluations of one of the (not many) permissible LDFs over  $\text{Dom}_{\star}(L)$ .

For each pair of local-readers, L of  $\mathcal{R}$  and M from  $\mathcal{R}_L$ , the composition procedure generates a local-reader of  $\mathcal{R}'$ , denoted by  $L \circ M$ , as follows. Let G denote the active domain of L, and let  $G[x_1], \ldots, G[x_t]$  denote its active variables. To obtain  $L \circ M$  each variable  $G[x_i]$  in the evaluator or the localtest of L is replaced by the *i*'th evaluator of M, and then the local-test of Mis added in conjunction to the local-test of L (where the  $G[x_i]$ 's have been replaced).

### Properties of the composition procedure

We now analyze the properties of the composition procedure that are important for its application by the CR constructor – the time it takes, and the properties and parameters of the LDF-readers it generates. In the analysis we assume that the composition procedure is applied to LDF-readers where the the number of variables in each local-reader and the number of conjunctions in each local-test is bounded by a constant, since the CR constructor indeed applies it to such LDF-readers. Notice that under this assumption it is clear that the composition procedure generates LDF-readers where the number of variables in each local-reader and the number of conjunctions in each local-test is also bounded by a (different) constant.

**Time.** When applied to an LDF-reader  $\mathcal{R}$ , the composition procedure applies the SP constructor several times. Each call to the SP constructor takes time polynomial in  $|\mathcal{F}|^{d_{\star}(\mathcal{R})}$ , according to the definition of a constructor (note that this is polynomial the number of variables in each of the active domains of  $\mathcal{R}$ ). Since the number of calls to the SP constructor equals the number of local-readers in  $\mathcal{R}$ , it follows that overall the composition procedure takes time polynomial in the size of  $\mathcal{R}$ .

**Encoding-assignments.** When the composition procedure is applied to an LDF-reader  $\mathcal{R}$  that evaluates a tuple  $(x_1, \ldots, x_k)$  in a domain F, the resulting structure  $\mathcal{R}'$  has representation-variables and local-readers. To be a valid LDF-reader, we show that for every good assignment  $\mathcal{A}$  for F there is an encoding-assignment with respect to  $\mathcal{R}'$ : First extend  $\mathcal{A}$  to an encodingassignment  $\mathcal{A}'$  for  $\mathcal{R}$ . In particular  $\mathcal{A}'$  assigns a good assignment to the active domain  $\text{Dom}_*(L)$  of each local-reader L in  $\mathcal{R}$ . Then extend the assignment of each active domain  $\text{Dom}_*(L)$  to an encoding-assignment with respect to  $\mathcal{R}_L$ . This obtains an assignment for all the variables of  $\mathcal{R}'$ . It is easy to verify that it is an encoding-assignment of  $\mathcal{A}$  with respect to  $\mathcal{R}'$ .

**Parameters of**  $\mathcal{R}'$ . Given an LDF-reader  $\mathcal{R}$ , the composition procedure generates an LDF-reader  $\mathcal{R}'$ . The parameters of  $\mathcal{R}'$  can be computed from the parameters of  $\mathcal{R}$  according to the following composition lemma.

**Lemma 4.15** (composition). Let  $\mathcal{R}$  be a restricted  $(\rho, \epsilon)$ -LDF-reader where  $\epsilon^{3/4} > (r_{\star}(\mathcal{R})/|\mathcal{F}|)^{c_g} d_{\star}(\mathcal{R})$ . Then the restricted LDF-reader  $\mathcal{R}'$ , generated from  $\mathcal{R}$  by the composition procedure, has parameters  $(\rho, O(\epsilon^{1/4}))$ .

Before the formal proof is given, we describe its main ideas. There are two types of  $\rho$ -erroneous local-readers  $L \circ M$ . One is where M is  $\epsilon^{3/4}$ -erroneous

– this happens for at most an  $O(\epsilon^{1/4})$  fraction of the local-readers since the  $\mathcal{R}_L$ 's are  $(\epsilon^{3/4}, O(\epsilon^{1/4}))$ -LDF-readers.

In case M is not erroneous, its evaluators yield evaluations of an  $\epsilon^{3/4}$ permissible LDF f with respect to the assignment of  $\text{Dom}_{\star}(L)$ .  $L \circ M$  is thus  $\rho$ -erroneous if and only if L remains  $\rho$ -erroneous when  $\text{Dom}_{\star}(L)$  is assigned
the feasible LDF f. Since for any active-feasible assignment for  $\mathcal{R}$  at most
an  $\epsilon$ -fraction of its local-readers may be  $\rho$ -erroneous, and since the number
of  $\epsilon^{3/4}$ -permissible LDFs for every domain is less than  $2\epsilon^{-3/4}$ , a counting
argument implies that the fraction of local-readers  $L \circ M$  where M is not
erroneous is bounded by  $2\epsilon^{-3/4} \cdot \epsilon = O(\epsilon^{1/4})$ .

## Proof of Lemma 4.15:

Fix an active-feasible assignment  $\mathcal{A}'$  for the representation of  $\mathcal{R}'$  and a parameter  $\rho' \doteq \epsilon^{3/4}$ , and let us divide the  $\rho$ -erroneous local-readers of  $\mathcal{R}'$ into two sets according to  $\rho'$  – the *peripheral-erroneous* local-readers are the local-readers  $L \circ M$  where M is  $\rho'$ -erroneous as a local-reader of  $\mathcal{R}_L$ , and the *core-erroneous* are those where M is not  $\rho'$ -erroneous. We bound the fraction of both types of local-readers by  $O(\epsilon^{1/4})$ .

**Peripheral-erroneous local-readers.** Since  $\rho' > (r_*(\mathcal{R})/|\mathcal{F}|)^{c_g} d_*(\mathcal{R})$ , Lemma 4.1 implies that every LDF-reader  $\mathcal{R}_L$  generated by the composition procedure has parameters  $(\rho', O((\rho')^{1/3}))$ , so the fraction of  $\rho'$ -erroneous local-readers in it is  $O((\rho')^{1/3}) = O(\epsilon^{1/4})$ . Hence for every local-reader L of  $\mathcal{R}$ , the fraction of peripheral-erroneous local-readers among local-readers of the form  $L \circ M$  is  $O(\epsilon^{1/4})$ , and therefore the overall fraction of peripheralerroneous local-readers in  $\mathcal{R}'$  is bounded by  $O(\epsilon^{1/4})$  as desired.

We move to bound the fraction of core-erroneous local-readers. We first show that in such a local-reader  $L \circ M$ , L has to be erroneous as a local-reader of  $\mathcal{R}$  with respect to a certain class of assignments, as explained below.

The assignments  $\mathcal{A}(G,g)$  for  $\mathcal{R}$ . For an active domain G of  $\mathcal{R}$  and an [r(G), d(G)]-LDF g, we define a class  $\mathcal{A}(G,g)$  of assignments for  $\mathcal{R}$ , based on  $\mathcal{A}'$ . The elements of  $\mathcal{A}(G,g)$  are the assignments for  $\mathcal{R}$  that assign g to G, and that are equal to  $\mathcal{A}'$  on all domains of  $\mathcal{R}$  which are not active. Active domains of  $\mathcal{R}$  other than G may be assigned arbitrarily. A local-reader L of  $\mathcal{R}$  with  $\text{Dom}_{\star}(L) = G$ , may be either  $\rho$ -erroneous with respect to all assignments in  $\mathcal{A}(G,g)$ , or with respect to none, because the assignments in

 $\mathcal{A}(G,g)$  are all equal on the variables of L (recall that L cannot have variables from active domains other than G).

Consider a local-reader  $L \circ M$  that is core-erroneous. The evaluators of M yield values consistent with an LDF g, which is  $\rho'$ -permissible with respect to the assignment of  $G \doteq \text{Dom}_{\star}(L)$ . It follows that as a local-reader of  $\mathcal{R}$ , L is  $\rho$ -erroneous with respect to the assignments in  $\mathcal{A}(G,g)$  – these assignments yield the same values for the variables of G as the evaluators of M, and give the same values as  $\mathcal{A}'$  to all the other variables of L.

**Core-erroneous local-readers.** Let G be an active domain of  $\mathcal{R}$ . Denote by  $\alpha(G, g)$  the fraction among  $\mathcal{R}$ -local-readers, of local-readers whose active domain is G and which are  $\rho$ -erroneous with respect to the assignments in  $\mathcal{A}(G, g)$ . Denote by  $\alpha(G)$  the maximum over all  $\alpha(G, g)$ .

Let  $\mathcal{A}$  be the assignment obtained from  $\mathcal{A}'$  by assigning to each active domain G of  $\mathcal{R}$  an LDF g such that  $\alpha(G, g)$  is maximized. Then for every G, the fraction of  $\mathcal{R}$ -local-readers whose active domain is G, and which are  $\rho$ -erroneous with respect to  $\mathcal{A}$  is  $\alpha(G)$ . The parameters of  $\mathcal{R}$  imply that the total fraction of  $\rho$ -erroneous local-readers is bounded by  $\epsilon$ , hence  $\sum_{G} \alpha(G) \leq \epsilon$ .

For an active domain G of  $\mathcal{R}$  we denote by  $\gamma(G)$  the fraction of localreaders L in  $\mathcal{R}$  whose active domain is G, and for which there exists a localreader M in  $\mathcal{R}_L$  where  $L \circ M$  is core-erroneous. We have seen that for an  $\mathcal{R}$ -local-reader L to be accounted in  $\gamma(G)$ , it must be  $\rho$ -erroneous with respect to the assignments in  $\mathcal{A}(G,g)$ , for some  $\rho'$ -permissible g.  $\gamma(G)$  is therefore bounded by the sum  $\sum \alpha(G,g)$  taken over all permissible LDFs g, and so by  $\alpha(G)$  times the number of  $\rho'$ -permissible LDFs. By Proposition 3.4 the number of  $\rho'$ -permissible LDFs is less than  $2/\rho'$ , hence  $\gamma(G) < (2/\rho')\alpha(G)$ , and we obtain that

$$\sum_{G} \gamma(G) < (2/\rho') \sum_{G} \alpha(G) \le 2\epsilon/\rho' = 2\epsilon^{1/4}$$

Namely the fraction of  $\mathcal{R}$ -local-readers L for which there exists a coreerroneous local-reader  $L \circ M$  is bounded by  $O(\epsilon^{1/4})$ .

We show that this also bounds the total fraction of core-erroneous localreaders: Note that there is the same number of local-readers in every LDFreader of the form  $\mathcal{R}_L$  – this follows from the definition of a constructor, together with the fact that all active domains of  $\mathcal{R}$  have the same degree parameters. Hence for each local-reader L of  $\mathcal{R}$  there is the same number of local-readers of the form  $L \circ M$  in  $\mathcal{R}'$ . A simple counting-argument now shows that the fraction of core-erroneous local-readers is also bounded by  $O(\epsilon^{1/4})$ , as desired.

# 4.6 The CR Constructor

It is now the time to describe the actual CR constructor, proving the Composition-Recursion Constructor Lemma (Lemma 3.2). Let F be a domain, and let  $(x_1, \ldots, x_k)$  be a k-tuple of points in  $\mathcal{F}^{d(F)}$  (where, as in Lemma 3.2, k is a constant). We assume, under the notation as specified in Lemma 3.2, that  $d(F) = O(\log^{1-\beta} n), \ s(F) \leq |\mathcal{F}|^{c_1}, \ \text{and} \ r(F) \geq |\mathcal{F}|^{c_2}$ . For simplicity we reset s(F) and r(F) so that the latter inequalities hold as equalities – note that a  $(\rho, \epsilon)$ -LDF-reader with respect to the new degree parameters remains a  $(\rho, \epsilon)$ -LDF-reader if s(F) is reduced and r(F) is increased to their original values.

The CR constructor generates an (unrestricted) LDF-reader  $\mathcal{R}$  evaluating  $(x_1,\ldots,x_k)$  in F. First, it generates a sequence  $\mathcal{R}_0,\ldots,\mathcal{R}_K$ , where  $K = O(\frac{1}{1-\beta})$  is a constant that will be chosen later, of restricted LDFreaders. The transformation of each  $\mathcal{R}_i$  into  $\mathcal{R}_{i+1}$  is accomplished in two steps. At first a restricted LDF-reader  $\mathcal{R}'_i$  is generated by applications of an extension-procedure to the active domains of  $\mathcal{R}_i$ , as described in the extension proposition (Proposition 4.12). The degree parameters of  $\mathcal{R}'_i$  are decreased with respect to  $\mathcal{R}_i$  but the dimension is increased.  $\mathcal{R}_{i+1}$  is then generated by applying the composition procedure to  $\mathcal{R}'_i$ , thus the degree parameters remains the same while the active dimension parameter becomes constant. Finally  $\mathcal{R}_K$  has a constant active dimension and both of its active degree parameters are 1, hence in a good or active-feasible assignment each active domain of  $\mathcal{R}_K$  is assigned a constant-dimensional linear function. The final unrestricted LDF-reader is obtained by replacing each active domain of  $\mathcal{R}_K$  by a constant number of variables that represent the coefficients of a linear function over it.

We now fully describe the generation of the sequence  $\mathcal{R}_0, \ldots, \mathcal{R}_K$ , and the transformation of  $\mathcal{R}_K$  into  $\mathcal{R}$ . We then show that the CR constructor has the properties required by Lemma 3.2.

**Generating**  $\mathcal{R}_0$ . To generate the first restricted LDF-reader,  $\mathcal{R}_0$ , the CR constructor applies the SP constructor to the domain F and the tuple  $(x_1, \ldots, x_k)$ .

**Generating**  $\mathcal{R}_1, \ldots, \mathcal{R}_{K-1}$ . From  $\mathcal{R}_0$  the CR constructor continues to iteratively generate restricted LDF-readers as follows. Having generated  $\mathcal{R}_i$ , the constructor transforms it into a restricted LDF-reader  $\mathcal{R}'_i$  by applying the power-substitution extension-procedure to each active domain of  $\mathcal{R}_i$  with parameter

$$b = \max\{(s_{\star}(\mathcal{R}_i) + 1)^{1/\log^{1-\beta} n}, 2\}$$

and taking these extensions to be the active domains of  $\mathcal{R}'_i$ . The constructor then generates  $\mathcal{R}_{i+1}$  by applying the composition procedure to  $\mathcal{R}'_i$ . The CR constructor iteratively generates LDF-readers as described above until finally an LDF-reader  $\mathcal{R}_{K-1}$  is generated such that

$$\binom{s_{\star}(\mathcal{R}_{K-1}) + d_{\star}(\mathcal{R}_{K-1})}{d_{\star}(\mathcal{R}_{K-1})} \leq \log^{1-\beta} n$$

As proven below, this occurs for a constant K.

**Generating**  $\mathcal{R}_K$ . The transformation of  $\mathcal{R}_{K-1}$  into  $\mathcal{R}_K$  is carried similarly to the previous transformations described above, only that  $\mathcal{R}'_{K-1}$  is generated using the linearization extension-procedure instead of the power-substitution extension-procedure. Note that for the linearization extension-procedure to be applicable the active lower-degree parameter of  $\mathcal{R}_{K-1}$  must not be greater than its active upper-degree. We show below that this indeed holds.

**Generating**  $\mathcal{R}$ . The constructor now transforms  $\mathcal{R}_K$  into the final CR LDF-reader. Having used the linearization extension-procedure to produce  $\mathcal{R}'_{K-1}$ , we gather that the active lower-degree of  $\mathcal{R}_K$  (which equals that of  $\mathcal{R}'_{K-1}$ ) is 1. Its active dimension,  $d \doteq d_*(\mathcal{R}_K)$ , is constant since  $\mathcal{R}_K$  is generated by the composition procedure. A good assignment to an active domain G of  $\mathcal{R}_K$  is thus a *linear* LDF f, that can be represented using a constant number of coefficient  $\gamma_i$  by

$$\forall (u_1, \dots, u_d) \in \mathcal{F}^d \qquad g(u_1, \dots, u_d) \doteq \sum_{i=1}^d \gamma_i u_i$$

The CR constructor adds d variables to the representation,  $G_1, \ldots, G_d$ , for each active domain G of  $\mathcal{R}_K$ , to represent the coefficients  $\gamma_1, \ldots, \gamma_d$  above. It then goes over all the local-readers and replaces every term  $G[(u_1, \ldots, u_d)]$ , where G is an active-LDF, by  $\sum_{i=1}^{d} G_i u_i$ . It is now possible to deactivate or even remove the active domains altogether (their variables no longer appear anywhere), thus completing the generation of  $\mathcal{R}$ .

# 4.7 The CR Constructor Works

Below it is proven that the CR constructor above has the properties stated in Lemma 3.2. We show that it stops after a constant number of iterations as stated above, and that it takes polynomial time. It is then shown that although each transformation of  $\mathcal{R}_i$  into  $\mathcal{R}_{i+1}$  consumes some of the lowerdegree to upper-degree gap, the active upper-degree of  $\mathcal{R}_{K-1}$  is not smaller than the active lower-degree (hence linearization extension-procedure is correctly used by the CR constructor). We conclude by showing that for an appropriate constant c > 0, the constructor generates  $(\rho, O(\rho^c))$ -LDF-readers for all  $\rho$ 's such that  $\rho > (r/|\mathcal{F}|)^{c_g} d$ .

First of all observe that as noted in the description of the properties of the composition procedure, for every constant i both the number of variables and the number of conjunctions in each local-reader are bounded by a constant.

## The number of iterations is constant.

It should be shown that for some constant  $K = O(\frac{1}{1-\beta})$ , the parameters of the (K-1)'th element in the sequence  $\mathcal{R}_0, \mathcal{R}_1, \ldots$  of LDF-readers satisfy

$$\binom{s_{\star}(\mathcal{R}_{K-1}) + d_{\star}(\mathcal{R}_{K-1})}{d_{\star}(\mathcal{R}_{K-1})} \le \log^{1-\beta} n \tag{2}$$

To see this we examine the parameters of the LDF-readers in the sequence.

**Parameters of the**  $\mathcal{R}_i$ 's. Consider an LDF-reader  $\mathcal{R}_i$  in the sequence, and assume  $s_{\star}(\mathcal{R}_i) + 1 > 2^{\log^{1-\beta} n}$ . The power-substitution extension-procedure is applied to its active domains using the parameter  $b = (s_{\star}(\mathcal{R}_i) + 1)^{1/\log^{1-\beta} n}$ , hence the parameter t used within the extension-procedure is  $t = \log^{1-\beta} n$  (see Proposition 4.13). Since the active dimension of  $\mathcal{R}_i$  is constant, Proposition 4.13 implies that

$$s_{\star}(\mathcal{R}_{i+1}) = s_{\star}(\mathcal{R}'_i) = d_{\star}(\mathcal{R}_i)t(b-1) = \operatorname{polylog}(n)s_{\star}(\mathcal{R}_i)^{1/\log^{1-\beta}n} \quad (3)$$

As  $\mathcal{R}_0$  is generated by the SP constructor, its active lower-degree parameter equals s(F), so  $s_{\star}(\mathcal{R}_0) = |\mathcal{F}|^{c_1} = 2^{\Theta(\log^{\beta} n)}$ . By inductively using Equation 3 one easily sees that as long as  $\beta - i(1 - \beta) > 0$ ,

$$s_{\star}(\mathcal{R}_i) = 2^{\Theta(\log^{\beta - i(1-\beta)} n)} \tag{4}$$

(the poly-logarithmic factor is absorbed in the exponent).

**Parameters of**  $\mathcal{R}_{i_o}$ . Fix  $i_o \doteq |\beta/(1-\beta)|$ , and note that it is constant (it depends only on  $\beta$ , which remains constant throughout the proof). We have

$$1 - \beta \ge \beta - (i_o - 1)(1 - \beta) > 0$$

hence by Equation 4,

$$s_{\star}(\mathcal{R}_{i_o-1}) = 2^{\Theta(\log^{\beta - (i_o-1)(1-\beta)} n)}$$

 $\mathcal{R}'_{i_o-1}$  is generated by applying the extension-procedure with parameter

$$b = \max\left\{ \left( 2^{\Theta(\log^{\beta - i_o(1-\beta)} n)} \right), \ 2 \right\} = O(1)$$

since  $\beta - i_o(1 - \beta) \leq 0$ . The parameter t used is poly-logarithmic in n, specifically  $t = O(\log^{\beta - (i_o - 1)(1 - \beta)} n) \leq O(\log^{1 - \beta} n)$ . It hence follows from Proposition 4.13 that  $s_{\star}(\mathcal{R}_{i_o}) = s_{\star}(\mathcal{R}'_{i_o - 1})$  is poly-logarithmic in n.

**Parameters of**  $\mathcal{R}_{i_o+1}$ . The power-substitution extension-procedure is applied with parameter b = 2 to generate  $\mathcal{R}'_{i_o}$  from  $\mathcal{R}_{i_o}$ . Since  $s_{\star}(\mathcal{R}_{i_o})$  is polylogarithmic, t is poly-log-logarithmic in n, and therefore  $s_{\star}(\mathcal{R}'_{i_o}) = s_{\star}(\mathcal{R}_{i_o+1})$  is also poly-log-logarithmic in n. Since  $d_{\star}(\mathcal{R}_{i_o+1})$  is constant, it follows that

$$\binom{s_{\star}(\mathcal{R}_{i_o+1}) + d_{\star}(\mathcal{R}_{i_o+1})}{d_{\star}(\mathcal{R}_{i_o+1})} \le \log^{1-\beta} n \tag{5}$$

Setting  $K \doteq i_o + 2$ , we have that  $K = O(1/(1-\beta))$  is constant and that Inequality 2 clearly holds for  $\mathcal{R}_{K-1} = \mathcal{R}_{i_o+1}$ .

## The lower – upper-degree gap remains.

Going from  $\mathcal{R}_{K-1}$  to  $\mathcal{R}'_{K-1}$ , the CR constructor applies the linearization extension-procedure to each active domain of  $\mathcal{R}_{K-1}$ . This procedure is only applicable to domains where the lower-degree is not greater than the upperdegree, hence we must show that  $s_{\star}(\mathcal{R}_{K-1}) \leq r_{\star}(\mathcal{R}_{K-1})$ .

Let us compute how  $s_{\star}(\mathcal{R}_{i+1})/r_{\star}(\mathcal{R}_{i+1})$  behaves with respect to  $s_{\star}(\mathcal{R}_i)/r_{\star}(\mathcal{R}_i)$ for  $0 \leq i < K - 1$ . Let  $b_i$  denote the *b*-parameter with which the powersubstitution extension-procedure is applied to the active domains of  $\mathcal{R}_i$  to obtain  $\mathcal{R}'_i$ , and let  $t_i$  denote the associated *t*-parameter. According to Proposition 4.13,

$$\frac{s_{\star}(\mathcal{R}_{i+1})}{r_{\star}(\mathcal{R}_{i+1})} = \frac{s_{\star}(\mathcal{R}'_i)}{r_{\star}(\mathcal{R}'_i)} \le \frac{s_{\star}(\mathcal{R}_i) \cdot d_{\star}(\mathcal{R}_i)t_i(b_i - 1)}{r_{\star}(\mathcal{R}_i)} = O\left(\frac{s_{\star}(\mathcal{R}_i)}{r_{\star}(\mathcal{R}_i)} \cdot t_i b_i\right)$$

hence the ratio between the active lower-degree and the active upper-degree is consumed by a factor of up to  $O(t_i b_i)$  in the transition from  $\mathcal{R}_i$  to  $\mathcal{R}_{i+1}$ .

Let us bound  $t_i b_i$ . By the choice of the parameters  $b_i$  it follows that  $t_i \leq \log^{1-\beta} n$  for all *i*. According to the above computations of  $s_{\star}(\mathcal{R}_i)$ , for  $1 \leq i < K-3$ 

$$b_i = \left(s_{\star}(\mathcal{R}_i) + 1\right)^{1/\log^{1-\beta} n} = \left(2^{\Theta(\log^{\beta - i(1-\beta)} n)}\right)^{1/\log^{1-\beta} n} = 2^{\Theta(\log^{\beta - (i+1)(1-\beta)} n)} = 2^{O(\log^{\beta - (1-\beta)} n)}$$

and therefore  $t_i b_i = 2^{O(\log^{\beta - (1-\beta)} n)}$ . For i = K - 3 or i = K - 2,  $b_i = O(1)$  so in these cases  $t_i b_i$  is poly-logarithmic in n.

The initial lower-degree upper-degree fraction is  $s_{\star}(\mathcal{R}_0)/r_{\star}(\mathcal{R}_0) = |\mathcal{F}|^{c_1-c_2} = 2^{-\Theta(\log^{\beta} n)}$ . This fraction is consumed in each of the constant number of iterations by either  $2^{O(\log^{\beta-(1-\beta)} n)}$  or a poly-logarithm, hence  $s_{\star}(\mathcal{R}_{K-1})/r_{\star}(\mathcal{R}_{K-1}) = 2^{-\Theta(\log^{\beta} n)}$  and in particular  $s_{\star}(\mathcal{R}_{K-1}) < r_{\star}(\mathcal{R}_{K-1})$ , as desired.

#### The CR constructor takes polynomial time

We need to show that the CR constructor takes polynomial time in  $|\mathcal{F}|^{d(F)}$ . Since

$$|\mathcal{F}|^{d(F)} = \left(2^{\log^{\beta} n}\right)^{\Theta(\log^{1-\beta} n)} = n^{\Theta(1)}$$

this is equivalent to showing that it takes polynomial time in n. The proof is by showing that the generation of each LDF-reader  $\mathcal{R}_i$  in the sequence  $\mathcal{R}_0, \mathcal{R}'_0, \mathcal{R}_1, \mathcal{R}'_1, \ldots, \mathcal{R}_K$  takes polynomial time in n and in the size of the predecessor of  $\mathcal{R}_i$  (clearly the time it takes to generate the final LDF-reader from  $\mathcal{R}_K$  is polynomial in the size of  $\mathcal{R}_K$ ). This implies that the CR constructor indeed takes polynomial time.

**Generating**  $\mathcal{R}_0$ . The CR constructor generates  $\mathcal{R}_0$  using the SP constructor, which does take time polynomial in  $|\mathcal{F}|^{d(F)}$ .

**Generating**  $\mathcal{R}_i$  for  $0 < i \leq K$ . The CR constructor generates  $\mathcal{R}_i$  by applying the composition procedure to  $\mathcal{R}'_{i-1}$ . As mentioned in Subsection 4.5, this takes polynomial time in the size of  $\mathcal{R}'_{i-1}$ .

Generating  $\mathcal{R}'_i$  for i < K - 1. The CR constructor generates  $\mathcal{R}'_i$  by applying the power-substitution extension-procedure to all active domains of  $\mathcal{R}_i$ . The time it takes is bounded by the size of  $\mathcal{R}_i$  times the time needed for each application of the extension-procedure. By the definition of extension-procedures, each such application takes polynomial time in  $|\mathcal{F}|^{d_{\star}(\mathcal{R}'_i)}$ . According to Proposition 4.13,  $d_{\star}(\mathcal{R}'_i) = d_{\star}(\mathcal{R}_i)t_i = O(t_i) \leq O(\log^{1-\beta} n)$  where  $t_i$  is as denoted in the degree-gap computation, hence  $|\mathcal{F}|^{d_{\star}(\mathcal{R}'_i)} = n^{O(1)}$ . Therefore each application of the extension procedure takes polynomial time in n, as needed.

**Generating**  $\mathcal{R}'_{K-1}$ . The difference between the generation of  $\mathcal{R}_K$  and that of the other  $\mathcal{R}'_i$ 's, is that the linearization extension-procedures is applied to each active domain instead of the power-substitution extension-procedure. Each such application still takes polynomial time in  $|\mathcal{F}|^{d_*(\mathcal{R}_{K-1})}$  but here  $d_*(\mathcal{R}_{K-1})$  is calculated according to Proposition 4.14,

$$d_{\star}(\mathcal{R}_{K-1}) = \begin{pmatrix} s_{\star}(\mathcal{R}_{K-1}) + d_{\star}(\mathcal{R}_{K-1}) \\ d_{\star}(\mathcal{R}_{K-1}) \end{pmatrix} \le \log^{1-\beta} n$$

 $|\mathcal{F}|^{d_{\star}(\mathcal{R}_{K-1})}$  is therefore still polynomial.

## $(\rho, \epsilon)$ -parameters of the CR constructor.

We now show that  $\mathcal{R}$  has parameters  $(\rho, O(\rho^{4^{-K}/3}))$  for all  $\rho$ 's that satisfy  $\rho > (r/|\mathcal{F}|)^{c_g} d$ . We first prove by induction that for all i,  $\mathcal{R}_i$  is a restricted

 $(\rho, O(\rho^{4^{-i/3}}))$ -LDF-reader: For  $\mathcal{R}_0$  it follows directly from Lemma 4.1. Assume now that  $\mathcal{R}_{i-1}$  is a restricted  $(\rho, O(\rho^{4^{-i+1/3}}))$ -LDF-reader. The extension proposition (Proposition 4.12) implies that  $\mathcal{R}'_{i-1}$  has the same parameters. Since  $\mathcal{R}_i$  is generated from  $\mathcal{R}'_{i-1}$  by the composition procedure, Lemma 4.15 yields that  $\mathcal{R}_i$  is a restricted  $(\rho, O(\rho^{4^{-i/3}}))$ -LDF-reader, as desired (the O notation here is justified since we only make a constant number of steps in the induction). Note that the requirement over  $\epsilon$  in Lemma 4.15 holds here, since we apply it with  $\epsilon^{3/4} = O(\rho^{4^{-i}}) \geq \rho > (r/|\mathcal{F}|)^{c_g} d$ .

By the above induction,  $\mathcal{R}_K$  is a restricted  $(\rho, O(\rho^c))$ -LDF-reader for  $c \doteq 4^{-K}/3$ . To show that  $\mathcal{R}$  has the same parameters, we define for each assignment  $\mathcal{A}$  of  $\mathcal{R}$  an *active-feasible* assignment  $\mathcal{A}'$  for  $\mathcal{R}_K$ , such that an  $\mathcal{R}_K$ -local-reader is  $\rho$ -erroneous with respect to  $\mathcal{A}'$  if and only if the  $\mathcal{R}$ -local-reader generated from it is  $\rho$ -erroneous with respect to  $\mathcal{A}$ . This would imply that  $\mathcal{R}$  has the same  $(\rho, \epsilon)$  parameters as  $\mathcal{R}_K$ .

 $\mathcal{A}'$  differs from  $\mathcal{A}$  only on active domains of  $\mathcal{R}_K$  – for an active domain G and a variable  $G[(u_1, \ldots, u_d)]$  in it we define

$$\mathcal{A}'(G[(u_1,\ldots,u_d)]) \doteq \sum_{i=1}^d \mathcal{A}(G_i)u_i$$

where the  $G_i$ 's are the new variables added in the generation of the final CR constructor.  $\mathcal{A}'$  assigns to each active domain G a linear LDF represented by the assignment of the  $G_i$ 's, and is hence active-feasible. It is clear from the construction of  $\mathcal{R}$  that a local-reader of  $\mathcal{R}_K$  is  $\rho$ -erroneous with respect to  $\mathcal{A}'$  if and only if the  $\mathcal{R}$ -local-reader obtained from it is  $\rho$ -erroneous with respect to  $\mathcal{A}$ .

# 5 The Sum-Check

In this section we prove the Sum-Check Lemma, Lemma 3.1. A reduction algorithm is shown that given a system  $\Psi$  of n quadratic-equations, where there are up to n variables in each equation, generates a system  $\Psi_{sc}$  whose variables belong to domains, and where every equation accesses only a constant number of variables. The reduction of  $\Psi$  into  $\Psi_{sc}$  is gap-preserving in the sense that if  $\Psi$  is completely satisfiable then  $\Psi_{sc}$  can be completely satisfiable by a **good** assignment to its domains; and if there is no assignment for  $\Psi$  that satisfies more than a  $\frac{2}{|\mathcal{F}|}$  fraction of its equations, then no **feasible**  assignment for  $\Psi_{sc}$  can satisfy more than a  $\frac{2}{|\mathcal{F}|}$  fraction of its equations as well.

The reduction begins with the given system  $\Psi_0 \doteq \Psi$ , and puts it through a constant number  $\left(O(\frac{1}{1-\beta})\right)$  of transformations, obtaining a sequence  $\Psi_0, \Psi_1, \ldots, \Psi_l$  of equation-systems. The final system  $\Psi_{sc}$  is generated by a small alteration of  $\Psi_l$ . The intermediate systems  $\Psi_i$  share a similar structure and are hence called *restricted equation-systems*, as defined below. However, the number of variables in their equations decreases from up to n in  $\Psi_0$ , to a constant in  $\Psi_l$ .

We continue as follows. The next subsection defines the structure of a restricted equation-system. The transformation of each restricted equation-system  $\Psi_i$  into  $\Psi_{i+1}$  (the transformation of  $\Psi_0$  into  $\Psi_1$  is an exception) is performed by an algorithm that is described in Subsection 5.2. This algorithm is used for the transformation of each intermediate system into the next, but it uses a different *representation-procedure* each time. Subsection 5.3 describes the properties of the different representation-procedures used, and of the algorithm which transforms  $\Psi_0$  into  $\Psi_1$ . The complete reduction of  $\Psi$  into  $\Psi_{sc}$  is finally described in Subsection 5.4.

The next subsections are dedicated to proving the correctness of the reduction (Subsection 5.5), and to the description and correctness proofs of the product-check and the representation-procedures used (Subsections 5.6, 5.7, 5.8, and 5.9).

# 5.1 Restricted Equation-Systems

All the systems  $\Psi_i$ , i = 1, 2, ..., l, in the sequence generated by the reduction algorithm have a similar structure. The following is an exact definition thereof.

**Definition 12** (restricted equation-systems). A restricted equation-system  $\Psi$ , is a quadratic equation-system where some domains are considered active. The dimension, and the upper and lower-degree parameters of the active domains must all be the same. These parameters are called the active dimension, active upper-degree and active lower-degree of  $\Psi$ , and are denoted by  $d_{\star}(\Psi)$ ,  $r_{\star}(\Psi)$  and  $s_{\star}(\Psi)$  respectively.

Each equation  $\psi \in \Psi$  is written in the form " $\psi_{\star} = \psi_{\mathbf{c}}$ ".  $\psi_{\star}$  is called the active part of  $\psi$  and  $\psi_{\mathbf{c}}$  is called the core of  $\psi$ . While  $\psi_{\mathbf{c}}$  may contain any

variable of  $\Psi$  and can have quadratic as well as linear terms,  $\psi_{\star}$  contains only linear terms and the variables in it must all be from one active domain, called the active domain of  $\psi$  and denoted by  $\text{Dom}_{\star}(\psi)$ . The variables that appear in  $\psi_{\star}$  are called the active variables of  $\psi$ .

The equations in all intermediate equation-systems will have only a constant number of variables in their core – all other variables appear in the active part of the equations. It is hence useful to denote the number of variables in the core of an equation and the number of active variables by different names.

**Definition 13** (active and core-dependency). Let  $\Psi$  be a restricted equationsystem. The active-dependency of an equation  $\psi \in \Psi$ , denoted by  $D_{\star}(\psi)$ , is defined as the number of variables in  $\psi_{\star}$ . The core-dependency of  $\psi$ ,  $D_{\mathbf{c}}(\psi)$ , is defined as the number of variables in  $\psi_{\mathbf{c}}$ . The active-dependency of  $\Psi$ , denoted by  $D_{\star}(\Psi)$ , is the maximum of  $D_{\star}(\psi)$  over all equations  $\psi \in \Psi$ . The core-dependency of  $\Psi$  is denoted  $D_{\mathbf{c}}(\Psi)$ , and is defined similarly.

As mentioned above, the core-dependency parameters of all the restricted equation-systems in the sequence  $\Psi_1, \ldots, \Psi_l$  are constants. The active-dependency parameter is decreased gradually until it becomes constant in  $\Psi_l$ . The total number of variables in an equation of  $\Psi_l$  is therefore constant, as required by Lemma 3.1 (this property is preserved in the final transition from  $\Psi_l$  to  $\Psi_{sc}$ ).

# 5.2 The Main Transformation-Scheme

The transformation of each restricted equation-system  $\Psi_i$  into the next is done by substituting each equation  $\psi$  of  $\Psi_i$  by a "representation" containing several new equations. The transformations of  $\Psi_i$  into  $\Psi_{i+1}$  where i = $1, \ldots, l-1$  are in fact of a more specific structure, and are carried out by the system-representation algorithm. An important part of this algorithm is the application of a representation-procedure to each equation in the system (a different representation-procedure is used for different transformations). We now define a representation-procedure, and then describe the systemrepresentation algorithm.

#### **Representation-procedures**

A representation-procedure is an algorithm that is applied to an equation  $\psi$ and produces a set  $\mathcal{E}_{\psi}$  of *conjunctions* of equations, and a new domain denoted by  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$  (the new conjunctions may have variables from  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$ ). The conjunctions of  $\mathcal{E}_{\psi}$  represent  $\psi$  in the sense that they are only satisfied by extensions to  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$ , of assignments that also satisfy  $\psi$  – feasible assignments which do not satisfy  $\psi$  will satisfy almost none of the conjunctions in  $\mathcal{E}_{\psi}$ .

For  $i = 1, \ldots, l$ ,  $\Psi_{i+1}$  is obtained from  $\Psi_i$  by applying a representationprocedure to each equation  $\psi \in \Psi_i$ , generating a system  $\Psi'_i$  of conjunctions which is the union of the sets  $\{\mathcal{E}_{\psi}\}_{\psi \in \Psi_i}$ .  $\Psi_{i+1}$  is then generated by replacing the conjunctions with equations as in Proposition 3.6. If the representationprocedure generates conjunctions with a small number of variables, then the dependency parameter of  $\Psi_{i+1}$  will be smaller than that of  $\Psi_i$  (eventually the dependency is constant). Also, the active-domains of  $\Psi_{i+1}$  are set to be the new domains generated by the representation-procedure, and hence the active parameters of  $\Psi_{i+1}$  are changed. Actually the reduction generates domains with different parameters, contrary to a requirement in Lemma 3.1. This is rectified by applying a simple technical method at the end of the reduction, that makes all the domains uniform.

An [s, d]-representation-procedure. An [s, d]-representation-procedure is an algorithm **A** that receives as input an equation  $\psi$  in a restricted equation-system  $\Psi$ , and generates a set  $\mathcal{E}_{\psi}$  of conjunctions of quadraticequations that "represent"  $\psi$ . It also generates a new domain denoted  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$  – the conjunctions in  $\mathcal{E}_{\psi}$  may have variables from  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$  in addition to any variables of  $\Psi$ . For a conjunction  $\chi \in \mathcal{E}_{\psi}$  we define the active domain of  $\chi$  to be  $\text{Dom}_{\star}(\chi) \doteq \text{Dom}_{\star}(\mathcal{E}_{\psi})$ . The variables of  $\chi$  that are associated with  $\text{Dom}_{\star}(\chi)$  are called the active variables of  $\chi$ .

The parameters  $r(\text{Dom}_{\star}(\psi))$ , s and d determine the parameters of the new domain, namely  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$  must satisfy  $r(\text{Dom}_{\star}(\mathcal{E}_{\psi})) = r(\text{Dom}_{\star}(\psi))$ ,  $s(\text{Dom}_{\star}(\mathcal{E}_{\psi})) = s$ , and  $d(\text{Dom}_{\star}(\mathcal{E}_{\psi})) = d$ . The running time of **A** should be polynomial in  $|\mathcal{F}|^d = |\text{Dom}_{\star}(\mathcal{E}_{\psi})|$  and the size of  $\psi$ .

**Extension and restriction properties.** For the conjunctions in  $\mathcal{E}_{\psi}$  to properly represent  $\psi$ , it is required that **A** has the following extension and restriction properties:

- Extension Property: For every good assignment  $\mathcal{A}$  for  $\Psi$  that satisfies  $\psi$  there should be an *s*-degree LDF such that if it is assigned to  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$ , all the conjunctions in  $\mathcal{E}_{\psi}$  are satisfied.
- Restriction Property: If a feasible assignment for  $\Psi$  and for  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$  satisfies at least an  $|\mathcal{F}|^{-1/2}$  fraction of the conjunctions in  $\mathcal{E}_{\psi}$ , then  $\psi$  is satisfied as well.

**Uniformity.** It is required that the parameters s and d be functions of  $\Psi$  alone, so that the parameters of  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$  are the same for all equations  $\psi \in \Psi$  to which  $\mathbf{A}$  is applied. The number of conjunctions in  $\mathcal{E}_{\psi}$  should also be independent of  $\psi$  (and be a function of  $\Psi$  alone). The number of equations in each conjunction of  $\mathcal{E}_{\psi}$  should all be the same, and they must be independent of  $\psi$  as well. In addition we require that the number of equations in each conjunction is bounded by O(d).

**Conjunction-structure.** The conjunctions of  $\mathcal{E}_{\psi}$  should have the following structure.  $\psi_{\mathbf{c}}$  may appear at most once in at most one equation of each conjunction  $\chi \in \mathcal{E}_{\psi}$ . Except for the terms in this copy of  $\psi_{\mathbf{c}}$ , all terms must be linear and the number of terms not associated with the domain  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$  must be bounded by a constant (that is, a number which is independent of  $\psi$  and  $\Psi$ ).

## The system-representation algorithm.

Let us now describe how a restricted equation-system  $\Psi_i$  is transformed into  $\Psi_{i+1}$  using a representation-procedure **A**.

- 1. First, **A** is applied to every equation  $\psi \in \Psi_i$ .
- 2. A system  $\Psi'_i$  of conjunctions is constructed by taking the union of the sets  $\{\mathcal{E}_{\psi}\}_{\psi \in \Psi_i}$ . Note that the number of equations in each conjunction of  $\Psi'_i$  is the same, and that each equation of  $\Psi_i$  results in the same number of conjunctions in  $\Psi'_i$ .
- 3.  $\Psi_{i+1}$  is generated by replacing each conjunction  $\chi \in \Psi'_i$  by all linearcombinations of its equations (with multiplicities, if the same equation occurs more than once). The active domain of each equation is set to be the same as that of the conjunction from which it originated. The

active variables of such an equation are thus the same as the active variables of the originating conjunction.

4. For each equation  $\xi \in \Psi_{i+1}$ ,  $\xi_*$  and  $\xi_c$  are defined as follows. The variables associated with  $\text{Dom}_*(\xi)$  are moved to the left-hand side of the equation, and the other variables to the right-hand side (it follows from the properties of **A** that variables associated with  $\text{Dom}_*(\xi)$  only appear in linear terms). The left-hand side of  $\xi$  is then defined to be the active part  $\xi_*$  of  $\xi$ , and the right-hand side is set to be its core,  $\xi_c$ .

Let us examine some of the properties of the system-representation algorithm.

The parameters of  $\Psi_{i+1}$ . Since the upper-degree parameter of the domains produced by the representation-procedure **A** are the same as the active upper-degree of  $\Psi_i$ , we have  $r_*(\Psi_{i+1}) = r_*(\Psi_i)$ . If **A** is an [s, d]-representation-procedure, then the active domains of  $\Psi_{i+1}$  will all have lower-degree s and dimension d, hence  $s_*(\Psi_{i+1}) = s$  and  $d_*(\Psi_{i+1}) = d$ .

**Time.** The system-representation algorithm takes polynomial time in the size of  $\Psi_i$  and  $|\mathcal{F}|^{d_{\star}(\Psi_{i+1})}$ . Especially note that step 3 is applicable in polynomial time in  $|\mathcal{F}|^{d_{\star}(\Psi_{i+1})}$  and in the size of  $\Psi_i$  – the uniformity property requires that the number of equations in each conjunction be bounded by  $O(d_{\star}(\Psi_{i+1}))$ , hence the number of equations produced for each conjunction is  $|\mathcal{F}|^{O(d_{\star}(\Psi_{i+1}))}$ .

**Core-dependency.** Note that the core-dependency of  $\Psi_{i+1}$  is larger by at most a constant than that of  $\Psi_i$ . Consider an equation  $\psi' \in \Psi_{i+1}$  whose origin is an equation  $\psi \in \Psi_i$ . It has the variables of  $\psi_c$ , and at most a constant number of variables not associated with  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$ . The other variables are associated with  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$ , and are hence active, so the core-dependency of  $\psi'$  is larger by only a constant than that of  $\psi$ .

The gap. The extension and restriction properties of the representationprocedure **A** that is used by the system-representation algorithm, ensure that the fraction of satisfiable equations in  $\Psi_i$  with respect to good or feasible assignments is close to the satisfiable fraction in  $\Psi_{i+1}$ . Here is a precise definition of this property. **Definition 14** (gap-preserving algorithm). An algorithm that transforms a given equation-system  $\Psi_i$  into an equation-system  $\Psi_{i+1}$  is said to be gap-preserving if it has the following properties:

- Completeness: If  $\Psi_i$  can be completely satisfied by a good assignment then so can  $\Psi_{i+1}$ .
- Soundness: If a feasible assignment for  $\Psi_{i+1}$  can satisfy a  $\gamma$ -fraction of its equations, then there exists a feasible assignment for  $\Psi_i$  satisfying at least a  $\gamma O(|\mathcal{F}|^{-1/2})$  fraction of its equations.

Note that even in a gap-preserving algorithm, the gap is actually consumed by an  $O(|\mathcal{F}|^{-1/2})$  fraction. The reduction deals with this by applying a simple gap amplification technique in the transformation from the system  $\Psi_l$  into the final system  $\Psi_{sc}$ .

### **Proposition 5.1.** The system-representation algorithm is gap-preserving.

*Proof.* Assume that the system-representation algorithm is applied to a restricted equation-system  $\Psi_i$  using a representation-procedure **A**, and outputs  $\Psi_{i+1}$ .

The completeness property is implied from the extension property of **A** as follows. Suppose  $\Psi_i$  is satisfied by an assignment  $\mathcal{A}$ . According to the extension property,  $\mathcal{A}$  can be extended to assign for each  $\psi$  an  $s(\text{Dom}_{\star}(\mathcal{E}_{\psi}))$  degree LDF to  $\text{Dom}_{\star}(\mathcal{E}_{\psi})$  such that the conjunctions in  $\mathcal{E}_{\psi}$  are satisfied. The system of conjunctions  $\Psi'_i$ , generated by the system-representation algorithm, is hence satisfied by this extended assignment, and therefore  $\Psi_{i+1}$  is satisfied as well by Proposition 3.6.

Let us now prove the soundness property. Assume that  $\Psi_{i+1}$  is  $\gamma$ -satisfiable by a feasible assignment  $\mathcal{A}$ . Then by Proposition 3.6,  $\Psi'_i$  is at least  $\gamma - |\mathcal{F}|^{-1}$ satisfied by  $\mathcal{A}$ . Recall that the number of conjunctions in the set  $\mathcal{E}_{\psi}$  is the same for every  $\psi \in \Psi_i$ . Hence for at least a  $\gamma - 2|\mathcal{F}|^{-1/2}$  fraction of the sets  $\mathcal{E}_{\psi}$ , a  $|\mathcal{F}|^{-1/2}$  fraction of the conjunctions are satisfied: Otherwise the fraction of satisfied conjunctions in  $\Psi_{i+1}$  would be less than  $\gamma - 2|\mathcal{F}|^{-1/2} + (1 - \gamma + 2|\mathcal{F}|^{-1/2})|\mathcal{F}|^{-1/2} < \gamma - |\mathcal{F}|^{-1}$ .

By the restriction property of  $\mathbf{A}$ , it follows that at least a  $\gamma - 2|\mathcal{F}|^{-1/2}$  fraction of the equations in  $\Psi_i$  are satisfied by  $\mathcal{A}$ . The proof is thus completed, noting that the restriction of  $\mathcal{A}$  to the variables of  $\Psi_i$  is feasible.

# 5.3 The Representation-Procedures

We now state the properties of the representation-procedures that are utilized in reducing  $\Psi$  to  $\Psi_{sc}$ . Only the properties that are needed for the reduction are discussed – the actual representation-procedures and the proofs of their stated properties appear later.

# Product-check.

The product-check algorithm is actually not a representation-procedure. Its properties are stated here since it is applied to  $\Psi_0$  to produce  $\Psi_1$ . The generated system  $\Psi_1$  is a restricted equation-system with just one domain, which is an active domain. Another important property of  $\Psi_1$  is that its equation are *condensed*, as defined below. This property is necessary since the arithmetization representation-procedure is applied to  $\Psi_1$  to obtain  $\Psi_2$ , and it can only be applied to systems with condensed equations. A more detailed explanation of the use of condensed equations appears in Subsection 5.7, which describes the arithmetization representation-procedure.

The principal cube of a domain. To define the principal cube we assume that for each non-negative number  $s < |\mathcal{F}|$  an arbitrary subset  $\mathcal{H}_s$  of  $\mathcal{F}$  is fixed, of size s + 1. The principal cube of a domain F is defined to be the subset  $(\mathcal{H}_{s(F)})^{d(F)} \subseteq \mathcal{F}^{d(F)}$ .

**Definition 15** (condensed equations). Let  $\psi$  be an equation or a conjunction with an active domain F, in a restricted equation-system.  $\psi$  is said to be condensed if all of its active variables are associated with points in the principal cube of F. A restricted equation-system where all of its equations are condensed is called condensed.

**Lemma 5.2** (product-check). Let  $\Psi$  be a system of *n* quadratic equations over  $\mathcal{F}$ , where there are at most *n* variables in each equation. There exists a gap-preserving polynomial time algorithm that given such a system, constructs a restricted equation-system  $\Psi_*$  that has the following properties:

- $D_{\mathbf{c}}(\Psi_*)$  is bounded by a constant.
- $\Psi_*$  has exactly one domain F which is the active domain of all of its equations. The parameters of F, that also determine the active degree

and dimension parameters of  $\Psi_*$ , are  $r(F) = |\mathcal{F}|^{1/4}$ ,  $s(F) = |\mathcal{F}|^{1/8}$ , and  $d(F) = \Theta(\log^{1-\beta} n)$ .

• The equations in  $\Psi_*$  are all condensed.

## Arithmetization.

The arithmetization representation-procedure uses a technique from [BFL91] to generate systems with a reduced active-dependency parameter. Given a condensed equation  $\psi$ , it produces a representation  $\mathcal{E}_{\psi}$  where the number of active variables in each conjunction is a function of the degree and dimension parameters of  $\text{Dom}_{\star}(\psi)$ . If these parameters are small enough, then the active-dependency is decreased. Note that the degree and dimension parameters themselves are not decreased, hence an iterative application of the arithmetization representation-procedure would not further reduce the dependency.

**Lemma 5.3** (arithmetization). Let  $\Psi$  be a restricted equation-system satisfying  $s_{\star}(\Psi)d_{\star}(\Psi) + r_{\star}(\Psi) < |\mathcal{F}|^{1/2}$  and  $s_{\star}(\Psi) > 2$ , and where all the equations are condensed.

There exists a  $[2d_{\star}(\Psi)s_{\star}(\Psi), d_{\star}(\Psi) + 1]$ -representation-procedure called **arithmetization**, applicable to the equations of such systems, that generates conjunctions with at most  $2d_{\star}(\Psi) \cdot s_{\star}(\Psi)$  active variables.

## Curve-extension.

When applied to equations with a small active-dependency parameter, the curve-extension representation-procedure generates domains with small degree and dimension parameters. The active dependency is not reduced (in fact it increases somewhat), but then the system-representation algorithm is applied to the resulting system using the arithmetization representationprocedure, and the decrease in the degree and dimension parameters is utilized to reduce the active dependency as well. By applying the systemrepresentation-procedures alternately, the reduction gradually reduces the active-dependency, the active degree and the dimension parameters of the intermediate systems.

**Lemma 5.4** (curve-extension). Let  $\Psi$  be a restricted equation-system such that  $s_{\star}(\Psi)D_{\star}(\Psi)$  and  $r_{\star}(\Psi)D_{\star}(\Psi)$  are smaller than  $|\mathcal{F}|^{1/2}$ . There exists

an [s, d]-representation-procedure called **curve-extension**, applicable to the equations of such systems, for

$$d \doteq \min \left\{ d_{\star}(\Psi), \log_2 \left( s_{\star}(\Psi) \cdot \mathcal{D}_{\star}(\Psi) \right) \right\}$$
  
and  $s \doteq d \cdot \max \left\{ \left( s_{\star}(\Psi) \cdot \mathcal{D}_{\star}(\Psi) \right)^{\frac{1}{d_{\star}(\Psi)}}, 2 \right\}$ 

that generates only condensed conjunctions.

## Linearization.

Applying the system-representation algorithm using the linearization representationprocedure obtains a system with constant active-dependency, as desired. However it is applicable in polynomial time only to systems where the active degree and dimension parameters are very small (the running time of a representation-procedure is polynomial in the size of the newly generated domains, which may become very large in the case of linearization). Hence the reduction generates a sequence of intermediate equation-systems where the active parameters are gradually reduced, until they finally become suitable for the linearization representation-procedure to be applied.

**Lemma 5.5** (linearization). Let  $\Psi$  be a system such that  $r_{\star}(\Psi)D_{\star}(\Psi)$  and  $s_{\star}(\Psi)D_{\star}(\Psi)$  are smaller than  $(|\mathcal{F}|^{1/2})/2$ .

There exists a  $[1, s_{\star}(\Psi)D_{\star}(\Psi)]$ -representation-procedure called linearization and is applicable to such systems, that generates conjunctions with at most 4 active variables.

Note that as mentioned above, for the linearization representation-procedure to be applicable within the reduction,  $s_{\star}(\Psi)D_{\star}(\Psi)$  should be in fact considerably smaller than the above bound of  $|\mathcal{F}|^{1/2}$ .

# 5.4 The Reduction Algorithm of $\Psi$ Into $\Psi_{sc}$

We now state the reduction algorithm that transforms  $\Psi$  into  $\Psi_{sc}$ , as claimed by Lemma 3.1. This algorithm is mostly a concatenation of the algorithms that were discussed above. Starting with  $\Psi = \Psi_0$ , the reduction algorithm applies the product-check algorithm to obtain  $\Psi_1$ , and from there it continues to use the system-representation algorithm, applying it a constant ( $O(\frac{1}{1-\beta})$ ) number of times with different representation-procedures. This yields a sequence of equation-systems  $\Psi_2, \ldots, \Psi_l$ .  $\Psi_{sc}$  is then obtained from  $\Psi_l$  by a simple transformation.

We next give the sequence of transformations and representation-procedures used to obtain  $\Psi_l$  from  $\Psi_0$ , and then describe how  $\Psi_{sc}$  is obtained from  $\Psi_l$ . In Subsection 5.5 it is shown that this reduction takes polynomial time in n, and that the generated system  $\Psi_{sc}$  has the desired properties. Subsection 5.5 also shows that although each representation-procedures is applicable only to systems with certain parameters, the reduction algorithm does use them correctly.

### The sequence of systems.

First, the reduction applies the product-check algorithm to  $\Psi_0$  and obtains  $\Psi_1$ . The next  $\frac{2\beta}{1-\beta}$  systems<sup>§</sup>,  $\Psi_2, \ldots, \Psi_{\frac{2\beta}{1-\beta}+1}$  are generated, by applying the system-representation algorithm with the arithmetization and the curve-extension representation-procedures alternately (the arithmetization is used first). The system-representation algorithm is then applied once more to  $\Psi_{\frac{2\beta}{1-\beta}+1}$  using the arithmetization representation-procedure, and then finally it is applied once again using the linearization representation-procedure. The outcome is  $\Psi_l$ , where  $l \doteq \frac{2\beta}{1-\beta} + 3$ . Apart from a simple transformation that is described shortly below,  $\Psi_l$  is the outcome of the reduction.

## Properties of $\Psi_l$ .

Before we describe how  $\Psi_l$  is transformed into  $\Psi_{sc}$ , let us overview its main properties.

**Constant dependency.**  $\Psi_l$  has the desired dependency parameter, namely a constant. Since it is generated using the linearization representationprocedure, it follows from Lemma 5.5 that its active-dependency parameter is constant. As for the core-dependency,  $\Psi_1$  is generated using the productcheck algorithm and therefore by Lemma 5.2 its core-dependency is constant. Since the other systems in the sequence  $\Psi_2, \ldots, \Psi_l$ , are generated using the system-representation algorithm, the core-dependency increases only by a

<sup>&</sup>lt;sup>§</sup>For simplicity of exposition, we assume here that  $\beta/(1-\beta)$  is an integer.

constant throughout the sequence (recall that the sequence is of constant length).

**Completeness, and soundness.** Since each of the intermediate transformations that were applied so far are gap-preserving, it follows immediately that the transformation from  $\Psi = \Psi_0$  into  $\Psi_l$  is gap preserving as well. Hence  $\Psi_l$  has the following properties:

- Completeness: If  $\Psi$  can be completely satisfied by a good assignment then so can  $\Psi_l$ .
- Weakened Soundness: If  $\Psi$  is no more than  $\frac{2}{|\mathcal{F}|}$ -satisfiable then  $\Psi_l$  cannot be more than  $O(|\mathcal{F}|^{-1/2})$ -satisfied by a feasible assignment.

## From $\Psi_l$ to $\Psi_{sc}$ .

 $\Psi_l$  fails to comply with two requirements of Lemma 3.1: The parameters of its domains are not all the same, and it has only a weakened soundness property, which is less than what is required in Lemma 3.1. The reduction hence transforms  $\Psi_l$  into  $\Psi_{sc}$  in two steps. First it resets the degree and dimension parameters of its domains without changing any of the other properties, and then it applies a simple technique to amplify the soundness property.

**Parameter uniformization.** First note that the upper-degree parameter is the same for all the domains of  $\Psi_l$  since  $\Psi_1$  has only one domain, and the representation-procedures generate domains with the same upper-degree as the active domain of the equation to which they are applied. Denote this upper-degree by  $r(\Psi_{sc})$ , and fix  $s(\Psi_{sc})$  to be the maximum over all lower-degrees of domains in  $\Psi_l$ , and  $d(\Psi_{sc})$  to be the maximum over all the dimension parameters. As shown in Subsection 5.5,  $s(\Psi_{sc})$  is smaller than  $r(\Psi_{sc})$ .

The reduction replaces each domain F of  $\Psi_l$  by a new domain F' with  $r(F') = r(\Psi_{sc}), s(F') = s(\Psi_{sc})$  and  $d(F') = d(\Psi_{sc})$ . Each variable F[x] which appears in an equation of  $\Psi_l$  is then replaced by the variable F'[x'], where x' is obtained from x by padding it with the appropriate number of zeros (in case the dimension parameter of F' is larger than that of F).

Note that the completeness and weakened soundness properties of  $\Psi_l$  are not affected by the uniformization step. Resetting the lower-degree parameter

maintains the completeness property since the lower-degree parameters may only be increased, and it has no effect on the soundness. The dimension enlargement also preserves the completeness property, as an LDF that was assigned to a domain before the change of dimension extends naturally to the larger domain maintaining the same degree, and thus a satisfying assignment can be translated through the uniformization step. Similarly, a feasible assignment to a domain with an enlarged dimension translates to a feasible assignment to the original domain by restriction, thus preserving the values of the variables appearing in the equations, and therefore the weakened soundness property is also maintained.

**Soundness amplification.** To amplify the soundness of  $\Psi_l$  the reduction first generates all conjunctions of three (not necessarily distinct) equations from  $\Psi_l$ . It then replaces each such conjunction with the set of all linear-combinations over its equations. The set of equations of  $\Psi_{sc}$  is thus

$$\{ \sum_{i=1}^{3} \lambda_i \psi_i : \forall i \quad \lambda_i \in \mathcal{F}, \ \psi_i \in \Psi_l \}$$

Completeness and soundness for  $\Psi_{sc}$ . Since it is simple to observe that the completeness property is maintained by the soundness amplification step, let us verify that  $\Psi_{sc}$  has the soundness property. Assume then that  $\Psi$  is no more than  $2/|\mathcal{F}|$ -satisfiable. As mentioned above, a feasible assignment for  $\Psi_l$  cannot satisfy more than an  $O(|\mathcal{F}|^{-1/2}) < |\mathcal{F}|^{-1/3}$  fraction of its equations, and this remains true when the domain-parameters of  $\Psi_l$  are reset. The fraction of conjunctions of three equations that can be satisfied by a feasible assignment is hence less than  $(|\mathcal{F}|^{-1/3})^3 = 1/|\mathcal{F}|$ . It then follows from Proposition 3.6 that  $\Psi_{sc}$  cannot be more than  $2/|\mathcal{F}|$ -satisfiable by a feasible assignment (Proposition 3.6 discusses general assignments but it is easily extendible to feasible assignments).

# 5.5 The Reduction Works

Based on the stated properties of the representation-procedures, we now verify that the reduction algorithm described above is applicable, and that the generated system  $\Psi_{sc}$  has the required parameters. The completeness and soundness properties of  $\Psi_{sc}$  have already been verified. From the properties of  $\Psi_l$  and the construction of  $\Psi_{sc}$  it is obvious that the number of variables in the equations of  $\Psi_{sc}$  is bounded by a constant and that the parameters of its domains are all the same.

We now compute the active parameters of all the intermediate systems  $\Psi_1, \ldots, \Psi_l$ , and at the same time verify that all representation-procedures are correctly used by the reduction. The computation will also imply that the parameters of the domains of  $\Psi_{sc}$  are as required by Lemma 3.1, and that the reduction takes polynomial time. For simplicity, we use O and  $\Theta$  notations in the computation, where any function that depends solely on  $\beta$  is regarded as constant.

#### The active parameters of the intermediate systems

As mentioned above the domains of  $\Psi_{sc}$ , as well as the domains in all the intermediate systems, all have the same upper-degree parameter, namely  $r(\Psi_{sc})$ . It also equals the active upper-degree of  $\Psi_1$ , hence  $r(\Psi_{sc}) = |\mathcal{F}|^{1/4}$ . Let us consider the other parameters of the intermediate systems.

The parameters of  $\Psi_1$ .  $\Psi_1$  is generated from  $\Psi_0$  using the product-check algorithm (see Lemma 5.2), hence it has the parameters

- $s_{\star}(\Psi_1) = |\mathcal{F}|^{1/8}$
- $d_{\star}(\Psi_1) = \Theta(\log^{1-\beta} n)$

The parameters of  $\Psi_2$ .  $\Psi_2$  is obtained from  $\Psi_1$  using the arithmetization representation-procedure. Note that the parameters of  $\Psi_1$  are such that the arithmetization representation-procedure is applicable. The parameters of  $\Psi_2$ , as follows from the arithmetization lemma, are

- $s_{\star}(\Psi_2) = 2d_{\star}(\Psi_1)s_{\star}(\Psi_1) = \Theta(|\mathcal{F}|^{1/8}\log^{1-\beta}n) = 2^{\Theta(\log^{\beta}n)}$
- $D_{\star}(\Psi_2) = 2d_{\star}(\Psi_1)s_{\star}(\Psi_1) = \Theta(|\mathcal{F}|^{1/8}\log^{1-\beta}n) = 2^{\Theta(\log^{\beta}n)}$
- $d_{\star}(\Psi_2) = d_{\star}(\Psi_1) + 1 = \Theta(\log^{1-\beta} n)$

The active parameters of  $\Psi_3, \Psi_4, \ldots, \Psi_{l-5}$  (recall that  $l-5 = \frac{2\beta}{1-\beta} - 2$ ) are given by the following proposition.

**Proposition 5.6.** For *i* such that  $3 \le 2i - 1 \le l - 6$ , the active parameters of  $\Psi_{2i-1}$  (generated using the curve-extension representation-procedure) are

- $s_{\star}(\Psi_{2i-1}) = 2^{\Theta(\log^{\beta i(1-\beta)} n)}$
- $d_{\star}(\Psi_{2i-1}) = \Theta(\log^{1-\beta} n)$

and for *i* such that  $4 \leq 2i \leq l-5$ , the parameters of  $\Psi_{2i}$  (that is generated using the arithmetization representation-procedure) are

- $s_{\star}(\Psi_{2i}) = 2^{\Theta(\log^{\beta i(1-\beta)} n)}$
- $D_{\star}(\Psi_{2i}) = 2^{\Theta(\log^{\beta i(1-\beta)} n)}$

• 
$$d_{\star}(\Psi_{2i}) = \Theta(\log^{1-\beta} n)$$

*Proof.* The proposition is obtained by induction over i, calculating the parameters of an equation-system according to the parameters of the previous system and the properties of the appropriate representation-procedure. We omit the calculation.

Note that the systems  $\Psi_{2i}$  have parameters such that the curve-extension representation-procedure is applicable, and that the arithmetization representationprocedure is applicable for the  $\Psi_{2i-1}$  systems, hence the sequence of transformation is valid up to and including  $\Psi_{l-5}$ . From the computations below it is also implied that the representation-procedure used for generating  $\Psi_{l-4}, \ldots, \Psi_l$  are also applicable.

**Parameters of**  $\Psi_{l-4}$ . Setting  $2i = l - 5 = 2(\frac{\beta}{1-\beta}) - 2$  in the above proposition we obtain that  $s_{\star}(\Psi_{l-5}) = D_{\star}(\Psi_{l-5}) = 2^{\Theta(\log^{1-\beta} n)}$ , and that  $d_{\star} = \Theta(\log^{1-\beta} n)$ . Hence according to the Curve-Extension Lemma (Lemma 5.4),

- $s_{\star}(\Psi_{l-4}) = \Theta(\log^{1-\beta} n) \cdot \Theta(1) = \Theta(\log^{1-\beta} n)$
- $d_{\star}(\Psi_{l-4}) = \Theta(\log^{1-\beta} n)$

**Parameters of**  $\Psi_{l-3}$ . The active parameters of this system, that is obtained using the arithmetization representation-procedure, are

- $s_{\star}(\Psi_{l-3}) = \Theta(\log^{2(1-\beta)} n)$
- $D_{\star}(\Psi_{l-3}) = \Theta(\log^{2(1-\beta)} n)$
- $d_{\star}(\Psi_{l-3}) = \Theta(\log^{1-\beta} n)$

**Parameters of**  $\Psi_{l-2}$ . The system  $\Psi_{l-2}$ , generated using the curve-extension representation-procedure, has parameters

- $s_{\star}(\Psi_{l-2}) = \Theta(\log \log n)$
- $d_{\star}(\Psi_{l-2}) = \Theta(\log \log n)$

**Parameters of**  $\Psi_{l-1}$ . This system, obtained via the arithmetization representationprocedure, is the last before the linearization representation-procedure is applied. Its parameters are

- $s_{\star}(\Psi_{l-1}) = \Theta(\log \log^2 n)$
- $D_{\star}(\Psi_{l-1}) = \Theta(\log \log^2 n)$
- $d_{\star}(\Psi_{l-1}) = \Theta(\log \log n)$

**Parameters of**  $\Psi_l$ .  $\Psi_{l-1}$  obviously satisfies the conditions of the Linearization Lemma (Lemma 5.5). According to the lemma, the parameters of  $\Psi_l$  are

- $s_{\star}(\Psi_l) = 1$
- $D_{\star}(\Psi_l) \leq 4$
- $d_{\star}(\Psi_l) = \Theta(\log \log^4 n)$

The parameters of  $\Psi_{sc}$ . By the above computations it is possible to deduce the parameters of the domains of  $\Psi_{sc}$ . Noting that  $s_{\star}(\Psi_2)$  is the highest active low-degree parameter of all intermediate systems it follows that  $s(\Psi_{sc}) = s_{\star}(\Psi_2) = \Theta(|\mathcal{F}|^{1/8} \log^{1-\beta} n)$ . Since  $r(\Psi_{sc}) = |\mathcal{F}|^{1/4}$ , it follows that the requirements over *s* and *r* in Lemma 3.1 hold. The above computations also imply that the active dimension of all intermediate systems is bounded by  $O(\log^{1-\beta} n)$ , and hence  $d(\Psi_{sc}) = \Theta(\log^{1-\beta} n)$  as required. **Polynomial time.** Since  $\Psi_{sc}$  was shown to satisfy all the requirements of Lemma 3.1, it is only left to verify that it is obtained from  $\Psi_0$  in polynomial time.  $\Psi_1$  is obtained in polynomial time, as stated in lemma 5.2. The other intermediate systems  $\Psi_2, \ldots, \Psi_l$ , are obtained by applying the system-representation algorithm. As stated in Subsection 5.2, an application of the system-representation algorithm to a system  $\Psi_{i-1}$  takes polynomial time in the size of  $\Psi_{i-1}$  and in  $|\mathcal{F}|^{d_*(\Psi_i)}$ .

According to the computations above  $d_{\star}(\Psi_i) = O(\log^{1-\beta} n)$  for all *i*, so  $|\mathcal{F}|^{d_{\star}(\Psi_i)}$  is polynomial in *n*. By induction it is therefore easy to verify that all intermediate systems are produced in polynomial time in *n*. The transformation of  $\Psi_l$  into  $\Psi_{sc}$  obviously takes polynomial time in the size of  $\Psi_l$ , so the entire reduction takes polynomial time in *n*.

# 5.6 The Product-Check Lemma

In this subsection we prove the product-check lemma. We show an algorithm that transforms a given quadratic-equation system into a restricted equationsystem with one domain, which has a relatively small (with respect to the size of the field) dimension parameter.

The product-check algorithm actually disposes of all the variables of  $\Psi$ , substituting them by the variables of the new domain F. Each variable of  $\Psi$  and each product of two such variables is replaced by a variables of the form F[x] that represent it. This is done so that for every assignment of  $\Psi$  there is a *good* assignment to F, where the value of each variable F[x] is equal to the value of the corresponding term in  $\Psi$ .

However, not every feasible assignment to F indeed represents an assignment of  $\Psi$ . Consider two variables of  $\Psi$  that are represented by  $F[x_1]$  and  $F[x_2]$  in F. There is no guarantee that the value of the variable F[x] that represents their product is indeed the product of the values of  $F[x_1]$  and  $F[x_2]$ . Each equation of  $\Psi$  is hence replicated several times in  $\Psi_*$ , where a "product-test" is added in conjunction to each copy to verify the correctness of the assignment.

## The product-check algorithm

Setting parameters and generating F. Let  $h \doteq |\mathcal{F}|^{1/9}$ , and let  $d \doteq \lceil \log_h(n+1) \rceil$ (note that  $d = O(\log^{1-\beta} n)$ ). The procedure constructs a new domain F with lower-degree parameter  $s(F) = |\mathcal{F}|^{1/8}$ , upper-degree  $r(F) = |\mathcal{F}|^{1/4}$ , and dimension d(F) = 2d.

**Representing terms.** The procedure chooses  $\mathcal{H} \subseteq \mathcal{H}_{s(F)} \subseteq \mathcal{F}$  to be an arbitrary set of size h. It then selects an arbitrary *injection*  $v \to x_v$ , associating every variable of  $\Psi$  with a point in  $\mathcal{H}^d \subseteq \mathcal{F}^d$  (such an injection exists). The procedure chooses another distinct point  $x_I \in \mathcal{H}^d$  to represent the value 1. Writing points in  $\mathcal{F}^{2d}$  as pairs  $(x_1, x_2)$  of points in  $\mathcal{F}^d$ , each variable v of  $\Psi$  is represented in  $\Psi_*$  by  $F[(x_v, x_I)]$ , and the product of two variables u, v is represented by  $F[(x_u, x_v)]$ .

**Generating conjunctions.** The procedure replaces each equation  $\psi$  of  $\Psi$  by a set  $\mathcal{E}_{\psi}$  of conjunctions as follows. Given  $\psi$ , it produces one conjunction in  $\mathcal{E}_{\psi}$  for every point  $(x_1, x_2) \in \mathcal{F}^{2d}$ , consisting of the following equations:

- 1.  $\psi$  itself, where every product  $u \cdot v$  is replaced by  $F[(x_u, x_v)]$  and every variable v in a linear term is replaced by  $F[(x_v, x_I)]$ .
- 2. The product-test equation  $F[(x_1, x_I)] \cdot F[(x_2, x_I)] = F[(x_1, x_2)].$
- 3. The equation  $F[(x_I, x_I)] = 1$ , which verifies that  $x_I$  indeed represents the value 1.

From conjunctions to equations. Let  $\Psi'$  denote the system of conjunctions, containing the union of all the sets  $\mathcal{E}_{\psi}$  where  $\psi \in \Psi$ . The system  $\Psi_*$  is generated from  $\Psi'$  by replacing each conjunction with all linear combinations of its equations, as described in Proposition 3.6. For every  $\chi \in \Psi_*$  we set  $\text{Dom}_{\star}(\chi)$  to be F.

Observing the construction of the conjunctions and of  $\Psi_*$ , one notes that there is at most one quadratic term in each equation  $\chi \in \Psi_*$ , and at most one more variable in each equation that is associated with a point outside  $\mathcal{H}^{2d}$ . These terms (the quadratic term and the additional variable), and also the constant term of each equation  $\chi$  are moved, if they exist, to the right-hand side of  $\psi$  and are set to be the core of  $\psi$ . The other terms are moved to the left-hand side, which is set to be the active part of  $\psi$ . The active variables of  $\psi$  are therefore all associated with points in  $\mathcal{H}^{2d}$ .

## **Proof of correctness**

It is easy to observe that the product-check algorithm indeed takes polynomial time. The generated system  $\Psi_*$  has one domain F, with parameters as stated by Lemma 5.2, and its core dependency is bounded by the constant 3 as required. Since the active variables of equations in  $\Psi_*$  are all associated with points in  $\mathcal{H}^{2d} \subseteq (\mathcal{H}_{s(F)})^{d(F)}$ , namely with points in the principal cube of F, we have that they are all condensed. It is left to show that the product-check algorithm is gap-preserving.

**Completeness.** Suppose  $\Psi$  is satisfiable by a good assignment  $\mathcal{A}$ . We show a good assignment  $\mathcal{A}'$  for F which represents it, namely that

- $\mathcal{A}'(F[(x_I, x_I)]) = 1$ , and for every variable v of  $\Psi$ ,  $\mathcal{A}'(F[(x_v, x_I)]) = \mathcal{A}(v)$ .
- $F[(x_1, x_I)] \cdot F[(x_2, x_I)] = F[(x_1, x_2)]$  for every  $x_1, x_2 \in \mathcal{F}^d$ .

It is easy to observe that an assignment  $\mathcal{A}'$  with the above properties will satisfy  $\Psi_*$ .

We define an LDF  $f : \mathcal{F}^d \to \mathcal{F}$  and then use it to define  $\mathcal{A}'$ . For points  $x_v \in \mathcal{H}^d$  associated with a variable v of  $\Psi$  we set  $f(x_v) \doteq \mathcal{A}(v)$ , and we also set  $f(x_I) \doteq 1$ . For points  $x \in \mathcal{H}^d$  not associated with variables, we arbitrarily set  $f(x) \doteq 0$ . We extend f over  $\mathcal{F}^d$  by the unique extension to an LDF of degree h - 1 in each variable. The total degree of f is therefore  $(h - 1)d = O(|\mathcal{F}|^{1/9} \log n)$ .  $\mathcal{A}'$  will assign to F the LDF g, defined by  $g(x_1, x_2) \doteq f(x_1) f(x_2)$ . This is a good assignment since g is of total-degree  $O(|\mathcal{F}|^{1/9} \log n) < |\mathcal{F}|^{1/8}$  (the inequality is true for large-enough n). The other stated properties of  $\mathcal{A}'$  are easy to verify.

**Soundness.** The next proposition is the first step in proving the soundness property. It shows that in order for  $\Psi_*$  to be  $|\mathcal{F}|^{-5/8}$ -satisfiable by a feasible assignment  $\mathcal{A}'$ ,  $\mathcal{A}'$  must be consistent with an assignment  $\mathcal{A}$  for  $\Psi$ . After proving the proposition we show that in that case  $\mathcal{A}$  must satisfy almost the same (up to  $\mathcal{F}^{-1}$ ) fraction of the equations in  $\Psi$  as  $\mathcal{A}'$  does for  $\Psi_*$ .

**Proposition 5.7.** Let  $\mathcal{A}'$  be an assignment of an r(F)-degree LDF g to F. If it satisfies at least an  $|\mathcal{F}|^{-5/8}$  fraction of the equations in  $\Psi_*$ , then there is an assignment  $\mathcal{A}$  for  $\Psi$  such that for every variable v of  $\Psi$ ,  $\mathcal{A}(v) = g(x_v, x_I)$ , and for every two variables u, v of  $\Psi \mathcal{A}(u)\mathcal{A}(v) = g(x_u, x_v)$ . *Proof.* Consider an assignment  $\mathcal{A}'$  as above, that assigns an LDF g to F and satisfies at least a  $|\mathcal{F}|^{-5/8}$  fraction of the equations of  $\Psi_*$ . We define an [r(F), d]-LDF f by  $f(x) \doteq g(x, x_I)$ , and set an assignment  $\mathcal{A}$  for every variable v of  $\Psi$  by  $\mathcal{A}(v) \doteq f(x_v)$  (hence the first stated property of  $\mathcal{A}$  holds).

By Proposition 3.6, if  $\mathcal{A}'$  satisfies more than an  $|\mathcal{F}|^{-5/8}$  fraction of the equations of  $\Psi_*$ , then it satisfies an  $\Omega(|\mathcal{F}|^{-5/8})$  fraction of the conjunctions in  $\Psi'$ . Then, for at least one of the equations  $\psi \in \Psi$ , the fraction of satisfied conjunctions in  $\mathcal{E}_{\psi}$  is at least  $\Omega(|\mathcal{F}|^{-5/8})$ . By observing the product-test in each conjunction of  $\mathcal{E}_{\psi}$ , we obtain that for an  $\Omega(|\mathcal{F}|^{-5/8})$  fraction of the points  $(x_1, x_2) \in \mathcal{F}^{2d}$ ,

$$f(x_1)f(x_2) = \mathcal{A}'(F[(x_1, x_I)])\mathcal{A}'(F[(x_2, x_I)]) = \mathcal{A}'(F[(x_1, x_2)]) = g(x_1, x_2)$$
(6)

In both sides of the equation we have LDFs of degree at most  $2r(F) = O(|\mathcal{F}|^{1/4})$ . Different LDFs of such parameters may only agree on an  $O(|\mathcal{F}|^{1/4}/|\mathcal{F}|) = O(|\mathcal{F}|^{-3/4})$  fraction of the points, however the LDFs in Equation 6 agree on an  $\Omega(|\mathcal{F}|^{-5/8})$  fraction and are hence equal. We therefore have

$$\forall (x_1, x_2) \in \mathcal{F}^{2d} \quad g(x_1, x_2) = f(x_1)f(x_2)$$

and specifically

$$\forall v, u \quad \mathcal{A}(u)\mathcal{A}(v) = f(x_u)f(x_v) = g(x_u, x_v)$$

as required.

We now return to the soundness proof of the product-check procedure. Assume that  $\Psi_*$  is  $\gamma$ -satisfiable by a feasible assignment  $\mathcal{A}'$ , and let us show an assignment  $\mathcal{A}$  satisfying a  $\gamma - O(|\mathcal{F}|^{-1/2})$  fraction of the equations in  $\Psi$ . We may assume that  $\gamma > |\mathcal{F}|^{-1/2}$  (otherwise there is nothing to show), and hence there exists an assignment  $\mathcal{A}$  for  $\Psi$  that corresponds to  $\mathcal{A}'$  as in Proposition 5.7.

The fraction of conjunctions in  $\Psi'$  that are satisfied by  $\mathcal{A}'$  is, by Proposition 3.6, at least  $\gamma - |\mathcal{F}|^{-1}$ . Hence for the same fraction of equations  $\psi$  of  $\Psi$ , there is at least one conjunction  $\chi \in \mathcal{E}_{\psi}$  which is satisfied by  $\mathcal{A}'$ . One of the equations in such a conjunction  $\chi$  is a copy of  $\psi$  where certain terms are replaced. According to Proposition 5.7 the replaced terms have the same value as the replacing terms, and therefore  $\psi$  is satisfied by  $\mathcal{A}$ . This implies that at least a  $\gamma - |\mathcal{F}|^{-1} > \gamma - |\mathcal{F}|^{-1/2}$  fraction of the equations of  $\Psi'$  are satisfied by  $\mathcal{A}$ .

## 5.7 The Arithmetization Representation-Procedure

In this subsection we show the arithmetization representation-procedure. When applied to an equation  $\psi$  whose active LDF has small degree and dimension parameters, this procedure produces a representation  $\mathcal{E}_{\psi}$  with small active-dependency. In essence, the arithmetization representation-procedure utilizes the *sum-check* technique, from previous PCP proofs (see [BFL91]).

We describe the running of the arithmetization representation-procedure over a given *condensed* equation  $\psi$  in a restricted equation-system  $\Psi$ . For shortness we denote  $E \doteq \text{Dom}_{\star}(\psi), r \doteq r(E), s \doteq s(E)$ , and  $d \doteq d(E)$ .

**Outline of the algorithm.** The equation  $\psi$  has the form  $\psi_{\star} = \psi_{\mathbf{c}}$ . Since  $\psi$  is condensed,  $\psi_{\star}$  can be written as a sum

$$\psi_{\star}$$
 :  $\sum_{y \in \mathcal{H}_s^d} \kappa(y) E[y]$ 

where  $\kappa : \mathcal{H}_s^d \to \mathcal{F}$  is a coefficient function. The assignment of the domain  $F \doteq \text{Dom}_{\star}(\mathcal{E}_{\psi})$ , that is generated by the arithmetization representationprocedure, encodes the summands in  $\psi_{\star}$  and also many of the partial sums. If the assignment of F is a correct encoding then it is possible to evaluate  $\psi_{\star}$ by accessing only a few variables of F, since their values are evaluations of large partial sums of  $\psi_{\star}$ .

Each conjunction of  $\mathcal{E}_{\psi}$  tests whether the assignment of F is a correct encoding, and assuming that the encoding is indeed correct, verifies that  $\psi$ holds. This is accomplished by accessing only a small number of variables of F. To satisfy more than an  $|\mathcal{F}|^{-1/2}$  fraction of the conjunctions in  $\mathcal{E}_{\psi}$  the assignment of F must encode the summands in the active part of  $\psi$  correctly, and also  $\psi$  must be satisfied.

The assignment for F that correctly encodes the summands of  $\psi$  is called the sum-check LDF, and is defined below. Values of the sum-check LDF at some points are evaluations of certain partial sums of  $\psi_{\star}$ , and values of the sum-check LDF at other points are used in testing the correctness of the encoding. It is necessary to define how the assignment of F should encode the summands in  $\psi_{\star}$  before the construction of  $\mathcal{E}_{\psi}$  can be understood, so we first describe the sum-check LDF and only then continue to the construction.

#### The sum-check LDF

We define the sum-check LDF of  $\psi$  with respect to a given good assignment  $\mathcal{A}$  for E. First, we extend  $\kappa$  to an LDF of degree ds over  $\mathcal{F}^d$  – such an extension exists and is computable in polynomial time in  $|\mathcal{F}|^d < |F|$ , and hence it is possible to compute the extension within the representation-procedure. We now define d LDFs that encode different partial sums of  $\psi_{\star}$ . The sum-check LDF is constructed from these LDFs below.

**Definition 16** (the sum-check tree). For k = 1, 2, ..., d, we define a function  $g_k : \mathcal{F}^k \to \mathcal{F}$  by

$$\forall x \in \mathcal{F}^k \quad g_k(x) \doteq \sum_{y \in \mathcal{H}_s^{(d-k)}} \kappa(x, y) \mathcal{A}(E[x, y])$$

where "x, y" means the concatenation of the vector x and the vector y. The sequence  $g_1, \ldots, g_d$  is called the sum-check tree with respect to  $\mathcal{A}(E)$ .

For an  $x \in \mathcal{H}_s^k$  the value of  $g_k(x)$  is a partial sum of  $\psi_{\star}$ . The value of  $g_d$ at a point  $x \in \mathcal{F}^d$  is just  $\kappa(x)\mathcal{A}(E[x])$ , and hence  $g_d$  is an LDF of degree at most ds + s = (d+1)s. It follows from the above definition that the other  $g_k$ 's have degree at most (d+1)s as well.

The LDFs  $g_1, \ldots, g_d$  form a tree of partial sums in the following sense. Consider a tree of depth d, where every non-leaf node has  $|\mathcal{F}|$  offsprings, and every node of depth k > 0 is labeled by a point evaluation of  $g_k$ . We label the root by  $\sum_{y \in \mathcal{H}_s^d} \kappa(y) \mathcal{A}(E[y])$ , which is the evaluation of  $\psi_{\star}$ . The root has an offspring labeled by  $g_1(z)$ , for each  $z \in \mathcal{F}$ . Note that for  $z \in \mathcal{H}_s$ ,  $g_1(z)$  is a partial sum of  $\psi_{\star}$ , and in fact the root-label is the sum of labels of its offsprings that are assigned  $g_1(z)$  for  $z \in \mathcal{H}_s$ .

For a non-leaf node that has been labeled  $g_k(x)$ , we label one of its offsprings by  $g_{k+1}(x, z)$  for every  $z \in \mathcal{F}$ . From the definition of the  $g_k$ 's it follows that for every k < d and  $x \in \mathcal{F}^k$ ,

$$g_k(x) = \sum_{z \in \mathcal{H}_s} g_{k+1}(x, z) \tag{7}$$

Hence the label of each node labeled  $g_k(x)$  in the tree is the sum of labels of its s + 1 offsprings that are assigned  $g_{k+1}(x, z)$  for  $z \in \mathcal{H}_s$ .

**The sum-check LDF.** We now incorporate all the LDFs  $g_1, \ldots, g_k$  into a single LDF of degree at most  $(d+1)s + d \leq 2ds$ , called the sum-check LDF. For this purpose, let  $\mathcal{H}_{d-1} = \{a_1, \ldots, a_d\}$  be an arbitrary subset of size d in  $\mathcal{F}$ . The sum-check LDF, denoted by f, will satisfy

$$f(a_k, x_1, \dots, x_k, 0, \dots, 0) = g_k(x_1, \dots, x_k)$$
(8)

for every  $1 \leq k \leq d$ , and every  $x = (x_1, \ldots, x_k) \in \mathcal{F}^k$ . There exists such an f – for example it can be defined by

$$f(x_0, x_1, \dots, x_d) \doteq \sum_{k=1}^d \left( \prod_{i \neq k} \frac{x_0 - a_i}{a_k - a_i} \right) \cdot g_k(x_1, \dots, x_k)$$

**Properties of the sum-check LDF.** From Equation 8 and the discussion above it follows that the sum-check LDF has the following properties:

- $\sum_{z \in \mathcal{H}_s} f[(a_1, z, 0, \dots, 0)]$  is the evaluation of  $\psi_*$ , as follows from the explanation after Definition 16.
- For  $k = 1, 2, \ldots, (d-1)$  and every  $(x_1, \ldots, x_k) \in \mathcal{F}^k$

$$f[(a_k, x_1, \dots, x_k, 0, \dots, 0)] = \sum_{z \in \mathcal{H}_s} f[(a_{k+1}, x_1, \dots, x_k, z, 0, \dots, 0)]$$

as follows from Equation 7.

• For every  $(x_1, \ldots, x_d) \in \mathcal{F}^d$ ,

$$f[(a_d, x_1, \dots, x_d)] = \kappa(x_1, \dots, x_d) \mathcal{A}(E[x_1, \dots, x_d])$$

as follows from the explanation after Definition 16.

#### The arithmetization representation-procedure

We now give the details of the arithmetization representation procedure. At first the representation-procedure produces a new domain  $F = \text{Dom}_{\star}(\mathcal{E}_{\psi})$ with parameters as stated in Lemma 5.3, namely r(F) = r, s(F) = 2dsand d(F) = d + 1. The procedure generates conjunctions that can only be satisfied if F is assigned the sum-check f. For each  $x = (x_1, \ldots, x_d) \in \mathcal{F}^d$ the procedure generates one conjunction, denoted by  $\chi[x]$ , consisting of the following d + 1 equations: • The root equation:

$$\sum_{z \in \mathcal{H}_s} F[a_1, z, 0, \dots, 0] = \psi_{\mathbf{c}}$$

• The d-1 path equations for  $k = 1, 2, \ldots, (d-1)$ :

$$F[a_k, x_1, \dots, x_k, 0, \dots, 0] = \sum_{z \in \mathcal{H}_z} F[a_{k+1}, x_1, \dots, x_k, z, 0, \dots, 0]$$

• The *leaf* equation:

$$F[a_d, x_1, \dots, x_d] = \kappa(x_1, \dots, x_d) E[x_1, \dots, x_d]$$

#### **Proof of correctness**

Let us show that the arithmetization representation-procedure has the required properties. It is easy to verify that it runs in polynomial time, and that it generates a domain  $F = \text{Dom}_{\star}(\psi)$  with parameters as required. As for the number of active variables in each conjunction, there are s + 1 variables associated with F in the root equation, s + 2 variables in each of the d - 1path equations, and one variable in the leaf equation. The total number is therefore  $s + 1 + (d - 1)(s + 2) + 1 = ds + 2d \leq 2ds$  as required. It is left only to verify the extension and restriction properties.

**Extension.** Let  $\mathcal{A}$  be a good assignment for the variables of  $\Psi$ . Extend  $\mathcal{A}$  to F by assigning the sum-check LDF f to it (f is of degree less than s(F)). From the properties of f stated above, it easily follows that if  $\psi$  is satisfied by  $\mathcal{A}$  then all the conjunctions of  $\mathcal{E}_{\psi}$  are also satisfied by the extension of  $\mathcal{A}$ .

**Restriction.** Let  $\mathcal{A}$  be a feasible assignment for the variables of  $\Psi$  and for F, and assume that at least an  $|\mathcal{F}|^{-1/2}$  fraction of the conjunctions in  $\mathcal{E}_{\psi}$  are satisfied. We define the sum-check tree  $g_1, \ldots, g_d$  and the sum-check LDF f with respect to the assignment of E, as in Definition 16 and Equation 8 above. Since now the degree of the LDF assigned to E may be up to r, the degree of the  $g_k$ 's can be up to  $sd + r < |\mathcal{F}|^{1/2}$ . We claim that F must be assigned f, at least at the points that matter, as stated in the following claim.
**Claim 5.8** (sum-check). Suppose that at least an  $|\mathcal{F}|^{-1/2}$  fraction of the conjunctions in  $\mathcal{E}_{\psi}$  are satisfied by a feasible assignment. Then for every k,  $1 \leq k \leq d$ , and every  $x = (x_1, \ldots, x_k) \in \mathcal{F}^k$ ,

$$\mathcal{A}(F[a_k, x_1, \dots, x_k, 0, \dots, 0]) = g_k(x_1, \dots, x_k)$$

Before proving the claim we show how it implies the restriction property. Note that the root equation is common to all the conjunctions in  $\mathcal{E}_{\psi}$ , and hence it must be satisfied. So together with the claim we have that the evaluation of  $\psi_{\mathbf{c}}$  equals  $\sum_{z \in \mathcal{H}_s} f(a_1, z, 0, \dots, 0)$ , which by the properties of the sum-check LDF equals the evaluation of  $\psi_{\star}$ . Therefore  $\psi$  is satisfied, as required.

**Proof of the sum-check claim.** For every  $k, 1 \le k \le d$ , we define an [r, k]-degree LDF  $g'_k$  by

$$g'_k(x_1,\ldots,x_k) \doteq \mathcal{A}(F[a_k,x_1,\ldots,x_k,0,\ldots,0])$$

For the sake of contradiction, assume that  $g'_k \neq g_k$  for some k, and choose k to be the highest for which this inequality holds. We distinguish between two cases for k:

- k = d: At least an  $|\mathcal{F}|^{-1/2}$  fraction of the conjunctions of  $\mathcal{E}_{\psi}$  are satisfied, and therefore at least the same fraction of the leaf equations are satisfied. So for at least an  $|\mathcal{F}|^{-1/2}$  fraction of the points  $x \in \mathcal{F}^d$ ,  $g'_d(x) = \mathcal{A}(F[a_d, x]) = \kappa(x)\mathcal{A}(E[x]) = g_d(x)$ . But according to our assumption  $g'_d \neq g_d$  and therefore their evaluations can not be equal on more than an  $\frac{sd+r}{|\mathcal{F}|} < |\mathcal{F}|^{-1/2}$  fraction of the points, a contradiction.
- $1 \leq k < d$ : At least an  $|\mathcal{F}|^{-1/2}$  fraction of the conjunctions of  $\mathcal{E}_{\psi}$  are satisfied, and therefore in at least the same fraction of them the k'th path equation is satisfied. It follows that for at least an  $|\mathcal{F}|^{-1/2}$  fraction of the points  $x = (x_1, \ldots, x_k) \in \mathcal{F}^k$ ,

$$g'_{k}(x) = \mathcal{A}(F[a_{k}, x_{1}, \dots, x_{k}, 0, \dots, 0]) =$$
  
=  $\sum_{z \in \mathcal{H}_{s}} \mathcal{A}(F[a_{k+1}, x_{1}, \dots, x_{k}, z, 0, \dots, 0]) =$   
=  $\sum_{z \in \mathcal{H}_{s}} g'_{k+1}(x_{1}, \dots, x_{k}, z)$ 

By the maximality of k we have that  $g'_{k+1} = g_{k+1}$ , hence for at least an  $|\mathcal{F}|^{-1/2}$  fraction of the points x,

$$g'_k(x) = \sum_{z \in \mathcal{H}_s} g_{k+1}(x_1, \dots, x_k, z) = g_k(x) \quad \text{(by Equation 7)}$$

This is a contradiction to our assumption that  $g'_k \neq g_k$ , since they are both of degree at most sd+r and therefore our assumption implies that they can be equal on at most an  $\frac{sd+r}{|\mathcal{F}|} < |\mathcal{F}|^{-1/2}$  fraction of the points.

# 5.8 The Curve-Extension Representation-Procedure

In this subsection we show the curve-extension representation-procedure. If it is applied to an equation with a small enough active-dependency, then the new generated domain has a small active lower-degree parameter, and for equations with even smaller active-dependency the active dimension parameter becomes small as well. The conjunctions that are generated by the procedure are all condensed.

Let us describe the running of the curve-extension representation-procedure over a given equation  $\psi$ . For shortness we denote  $E \doteq \text{Dom}_{\star}(\psi)$ ,  $r \doteq r(E)$ ,  $s \doteq s(E)$ ,  $d \doteq d(E)$ , and  $D \doteq D_{\star}(\psi)$ .

The principle of the algorithm. Denote the active variables of  $\psi$  by  $E[x_1], \ldots, E[x_D]$ . We define below a polynomial vector function of small degree  $\Gamma : \mathcal{F} \to \mathcal{F}^d$ , that goes through the points  $x_1, \ldots, x_D$ . The assignment of the domain  $F \doteq \text{Dom}_{\star}(\mathcal{E}_{\psi})$ , generated by the curve-extension representation-procedure, encodes the restriction of the assignment of E to the points of the curve  $\Gamma$ .

Variables in F associated with certain points in its principal cube have, in a correct encoding, the values of the assignment of E at certain points on  $\Gamma$ . The values at other points on  $\Gamma$  can be computed by interpolation over these variables of F, making use of the fact that  $\Gamma$  has a small degree, and hence restricting the assignment of E to its points yields an LDF of small degree as well. The conjunctions of  $\mathcal{E}_{\psi}$  use the variables of F to evaluate  $\psi_{\star}$ and verify that  $\psi$  is satisfied, and they also test whether F is indeed given a correct encoding.

#### The curve-extension algorithm

At first the representation-procedure produces a new domain  $F = \text{Dom}_{\star}(\mathcal{E}_{\psi})$ with parameters as stated in Lemma 5.4, that is

$$r(F) = r, \ d(F) = \min \{d, \log_2(sD)\}, \text{ and } s(F) = d(F) \cdot \max \{(s \cdot D)^{1/d}, 2\}$$

Each element of  $\mathcal{E}_{\psi}$  will be a conjunction of two condensed equations. One is an equation  $\psi'$ , derived from  $\psi$  by replacing each of its active variables with a variable of F that "encodes" it. The other equation is taken from a set of equations called a *curve-verifier*. These equations are not satisfied unless the assignment of F is a correct encoding. Before the construction of these equations, we define the curve  $\Gamma$  and describe how the assignment of F encodes the restriction of the assignment of E to the points of  $\Gamma$ .

**Definition 17** (the curve  $\Gamma$ ). Let  $\mathcal{H}_{sD-1}$  be an arbitrary subset of  $\mathcal{F}$  of size sD, and denote its elements by  $a_1, \ldots, a_{sD}$ .  $\Gamma : \mathcal{F} \to \mathcal{F}^d$  is defined to be the (D-1)-degree polynomial vector function satisfying

$$\forall \ 1 \le i \le D \qquad \Gamma(a_i) = x_i$$

where  $E[x_1], \ldots, E[x_D]$  are the active variables of  $\psi$ .  $\Gamma$  can clearly be computed in polynomial-time.

Associating points with  $a_1, \ldots, a_{sD}$ . Let  $(\mathcal{H}_{s(F)})^{d(F)}$  be the principal cube of F. The procedure chooses an arbitrary subset  $\mathcal{H} \subseteq \mathcal{H}_{s(F)}$  of size  $s(F)/d(F) = \max\{(s \cdot D)^{1/d}, 2\}$ , and associates to each point  $a_i$  in  $\mathcal{H}_{sD-1}$  a distinct point  $y_i$  in  $\mathcal{H}^{d(F)}$  (note that  $\mathcal{H}^{d(F)}$  is a subset of the principal cube of F and that it contains at least sD points). Each of the variables  $F[y_i]$  will encode the value of  $E[\Gamma(a_i)]$ . The active variables of the conjunctions in  $\mathcal{E}_{\psi}$ will all be of the form  $F[y_i]$ , so the conjunctions of  $\mathcal{E}_{\psi}$  are condensed. It is important to note that any assignment to the variables  $F[y_i]$  can be extended by interpolation to a good assignment for F, as is shown below.

**Generating the curve-verifier.** Suppose E is assigned an LDF g. Then a correct encoding assigns to  $F[y_i]$  the value of g at  $\Gamma(a_i)$ . Since  $\Gamma$  is of degree at most D - 1, if g is of degree s then  $g \circ \Gamma$  is of degree less than sD - 1. The value of g at any point on the curve  $\Gamma$  can hence be evaluated by interpolation over its values at  $\Gamma(a_1), \ldots, \Gamma(a_{sD})$  or, if F is assigned a correct encoding, by interpolation over the variables  $F[y_1], \ldots, F[y_{sD}]$ . This is stated precisely in the following claim.

**Claim 5.9** (curve-interpolation). Let s and D be such that  $sD < |\mathcal{F}|$ . Then there exists a polynomial (in  $|\mathcal{F}|$ ) algorithm that receives as input a point  $x \in \mathcal{F}$  and outputs a coefficient function  $\kappa_x : \{a_1, \ldots, a_{sD}\} \to \mathcal{F}$  with the following property: Every function  $f' : \{a_1, \ldots, a_{sD}\} \to \mathcal{F}$  has a unique extension to an [sD - 1, 1]-LDF f over  $\mathcal{F}$ , and f satisfies

$$\forall x \in \mathcal{F}$$
  $f(x) = \sum_{i=1}^{sD} \kappa_x(a_i) f'(a_i)$ 

The curve-verifier will have one equation  $\chi[x]$  for each point  $x \in \mathcal{F}$ .  $\chi[x]$  verifies that the interpolation over  $F[y_1], \ldots, F[y_{sD}]$  using the  $\kappa_x$  from Claim 5.9 yields the value of  $E[\Gamma(x)]$ , as it should if F is assigned the encoding of a good assignment to E:

$$\chi[x] : \sum_{i=1}^{sD} \kappa_x(a_i) F[y_i] = E[\Gamma(x)]$$

The next claim shows that the curve-verifier equations cannot be satisfied unless F is indeed assigned a correct encoding.

**Claim 5.10.** Let  $\mathcal{A}$  be a feasible assignment for E and F. Let f be the [sD - 1, 1]-LDF defined by  $f(x) = \sum_{i=1}^{sD} \kappa_x(a_i)\mathcal{A}(F[y_i])$ , as in Claim 5.9. Then either  $\mathcal{A}(E) \circ \Gamma = f$ , in which case all of the curve-verifier equations are satisfied, or less than an  $|\mathcal{F}|^{-1/2}$  fraction of the curve-verifier equations are satisfied.

Proof. Note that an equation  $\chi[x]$  of the curve-verifier is satisfied if and only if  $E[\Gamma(x)]$  is assigned f(x). It is thus obvious that these equations will all be satisfied if  $\mathcal{A}(E) \circ \Gamma = f$ . If this is not the case, then  $\mathcal{A}(E) \circ \Gamma$  and f are in particular two different [rD, 1]-LDFs. Since  $rD < |\mathcal{F}|^{1/2}$  it follows that their evaluations differ on all but less than an  $|\mathcal{F}|^{1/2}/|\mathcal{F}| \leq |\mathcal{F}|^{-1/2}$  fraction of the points. Hence if  $\mathcal{A}(E) \circ \Gamma \neq f$ , then less than an  $|\mathcal{F}|^{-1/2}$  fraction of the curve-verifier equations can be satisfied. **Generating**  $\psi'$ . The procedure generates an equation  $\psi'$  by replacing each active variable  $E[x_i]$  in  $\psi_*$  with the variable  $F[y_i]$ . If F is assigned a correct encoding then  $\psi'$  simulates  $\psi$ , as stated in the following claim.

**Claim 5.11.** Let  $\mathcal{A}$  be an assignment for  $\Psi$  and for F. Let f be the [sD - 1, 1]-degree LDF defined by  $f(x) = \sum_{i=1}^{sD} \kappa_x(a_i)\mathcal{A}(F[y_i])$ , as in Claim 5.9, and assume that  $\mathcal{A}(E) \circ \Gamma = f$ . In that case  $\psi$  is satisfied by  $\mathcal{A}$  if and only if  $\psi'$  is satisfied by it.

Proof. According to the definition of  $\Gamma$ ,  $\Gamma(a_i) = x_i$  for  $i = 1, \ldots, D$ . Hence it follows from the assumption that  $\mathcal{A}(E[x_i]) = \mathcal{A}(E[\Gamma(a_i)]) = f(a_i)$  for every  $i, 1 \leq i \leq D$ . But according to Claim 5.9  $f(a_i) = \mathcal{A}(F[y_i])$  for every i. Therefore the assignment of every active variable  $E[x_i]$  equals the assignment of  $F[y_i]$ . The claim immediately follows.

**Generating**  $\mathcal{E}_{\psi}$ . The set  $\mathcal{E}_{\psi}$  is composed of all the conjunctions of  $\psi'$  and an equation  $\chi[x]$  of the curve-verifier.

### **Proof of correctness**

The domain F that is generated by the curve-extension representation-procedure has the parameters required by Lemma 5.4, and the conjunctions of  $\mathcal{E}_{\psi}$  are all condensed. It is also easy to verify that the curve-extension representationprocedure takes polynomial time in the size of  $\psi$  and in |F|. To complete the proof of Lemma 5.4 it remains to show that  $\mathcal{E}_{\psi}$  has the extension and restriction properties. The other properties required of a representation-procedure are obvious.

• Extension: Let  $\mathcal{A}$  be a good assignment for the variables of  $\Psi$  that satisfies  $\psi$ . We extend  $\mathcal{A}$  by assigning an s(F)-degree LDF to F such that all the conjunctions of  $\mathcal{E}_{\psi}$  are satisfied. The LDF g, to be assigned to F, is defined as follows. First, let  $g(y_i) \doteq \mathcal{A}(E[\Gamma(a_i)])$  for  $i = 1, \ldots, sD$ . Since all the  $y_i$ 's are contained in  $\mathcal{H}^{d(F)}$ , there exists an extension of gto an LDF over  $\mathcal{F}^{d(F)}$  of degree at most s(F)/d(F) in each variable. The total degree of this g is hence at most s(F). We assign g to F. Then  $\mathcal{A}(F[y_i]) = \mathcal{A}(E[\Gamma(a_i)])$  for every i, and so Claim 5.11 implies that  $\psi'$  is satisfied. Let f be the [sD-1, 1]-LDF defined by  $f \doteq \mathcal{A}(E) \circ \Gamma$ . Then by Claim 5.9,

$$\forall x \in \mathcal{F} \qquad f(x) = \sum_{i=1}^{sD} \kappa_x(a_i) f(a_i) = \sum_{i=1}^{sD} \kappa_x(a_i) \mathcal{A}(E[\Gamma(a_i)])$$
$$= \sum_{i=1}^{sD} \kappa_x(a_i) \mathcal{A}(F[y_i])$$

where the coefficients  $\kappa_x(a_i)$  are as in Claim 5.9. It hence follows that the curve-verifier equations are all satisfied by the extended  $\mathcal{A}$ . Since  $\mathcal{E}_{\psi}$  consists of conjunctions of  $\psi'$  and equations of the curve-verifier, we have that all of its conjunctions are satisfied.

• Restriction: Let  $\mathcal{A}$  be a feasible assignment for the variables of  $\Psi$  and for F, and assume that at least an  $|\mathcal{F}|^{-1/2}$  fraction of the conjunctions in  $\mathcal{E}_{\psi}$  are satisfied by  $\mathcal{A}$ . Since  $\psi'$  appears in every conjunction of  $\mathcal{E}_{\psi}, \psi'$  is satisfied, and at least an  $|\mathcal{F}|^{-1/2}$  fraction of the curve-verifier equations are satisfied as well.

Define an (sD - 1)-degree LDF f by

$$\forall x \in \mathcal{F}$$
  $f(x) \doteq \sum_{i=1}^{sD} \kappa_x(a_i) \mathcal{A}(F[y_i])$ 

where the coefficients  $\kappa_x(a_i)$  are as in Claim 5.9. It follows from Claim 5.10 that  $\mathcal{A}(E) \circ \Gamma = f$ . Since  $\psi'$  is satisfied, it then follows from Claim 5.11 that  $\psi$  is satisfied as well, thereby proving the restriction property.

# 5.9 The Linearization Representation-Procedure

In this subsection we show the Linearization representation-procedure. It is the final representation-procedure used in the sequence of transformations, resulting in a system of a constant active-dependency parameter.

The linearization representation-procedure is similar to the curve-extension. When applied to an equation  $\psi$ , it uses the newly generated domain to encode the restriction of the assignment of  $\text{Dom}_{\star}(\psi)$  to a curve that contains the active variables of  $\psi$ . The curve-extension representation-procedure encoded directly the assignment at only some points of the curve; to obtain other evaluations it applied interpolation by computing an appropriate linearcombinations over the encoded values.

The linearization representation-procedure applies a method of [ALM<sup>+</sup>92], using the newly generated domain to encode all linear-combinations of these values. Hence each curve-verifier equation requires just one active variable of the new domain. Also, since the active part of  $\psi$  is a linear-combination of variables associated with points on the curve,  $\psi_{\star}$  can also be evaluated using one access to the new domain.

#### The linearization representation-procedure

We now describe the linearization representation-procedure. We fix the notations  $E \doteq \text{Dom}_{\star}(\psi)$ ,  $r \doteq r(E)$ ,  $s \doteq s(E)$ ,  $d \doteq d(E)$ , and  $D \doteq D_{\star}(\psi)$ . The linearization representation-procedure first generates a new domain F with parameters as stated in Lemma 5.5, that is

$$r(F) = r$$
,  $s(F) = 1$ , and  $d(F) = sD$ 

In each conjunction in  $\mathcal{E}_{\psi}$  there will be an equation  $\psi'$ , that is derived by replacing the active part of  $\psi$  with a variable of F that encodes it. Another equation in each conjunction is taken from a set of equations called a *linearization-verifier*, that are not satisfied unless F is assigned a homogeneous linear-LDF. As in the curve-extension representation-procedure, the last equation in each conjunction is taken from a set called the *curve-verifier*, whose equations are not satisfied unless the assignment of F is a correct encoding.

Generation of the linearization-verifier. The linearization-verifier has one equation  $\chi[y, t]$  for every  $y \in \mathcal{F}^{d(F)}$  and  $t \in \mathcal{F}$ :

$$\chi[y,t]$$
:  $tF[y] = F[ty]$ 

The equations of the linearization-verifier are not satisfied unless F is assigned a linear homogeneous LDF. To prove it we need the following observation.

**Claim 5.12.** Let f be an [r(F), d(F)]-LDF which is not linear homogeneous. For every point  $y \in \mathcal{F}^{sD}$  we define an [r(F), 1]-LDF  $\phi_y$  by

$$\forall t \in \mathcal{F} \qquad \phi_y(t) \doteq f(ty)$$

Then  $\phi_y$  is linear homogeneous for less than an  $(|\mathcal{F}|^{-1/2})/2$  fraction of the points y.

Proof. If f is linear but not homogeneous then obviously none of the  $\phi_y$ 's are homogeneous, so we assume that  $m \doteq \deg(f) > 1$ . Then f can be written as a sum  $f = f_1 + f_2$  where  $f_1$  is a homogeneous<sup>¶</sup> function of degree m, and  $\deg(f_2) < m$ . Then  $\phi_y(t) = f_1(y)t^m + f_2(ty)$ . The second term in this sum is of degree less than m as an LDF over t. The first term is an LDF of degree exactly m in t (and in particular it is not linear), unless  $f_1(y) = 0$ .

It follows that  $\phi_y$  is not linear (and in particular not linear-homogeneous) unless  $f_1(y) = 0$ , which occurs for at most an  $\frac{m}{|\mathcal{F}|} \leq \frac{r(F)}{|\mathcal{F}|} \leq |\mathcal{F}|^{-1/2}/2$  fraction of the points y, as we needed to show.

The next claim shows that the linearization-verifier equations are indeed satisfied only if F is assigned a linear-homogeneous function.

**Claim 5.13.** Let  $\mathcal{A}$  be a feasible assignment for F. Then less than an  $|\mathcal{F}|^{-1/2}$  fraction of the linearization-verifier equations are satisfied unless F is assigned a linear-homogeneous function, in which case all of the equations are satisfied.

Proof. It is clear that the linearization-verifier equations are all satisfied in the case that F is assigned a linear-homogeneous function. We therefore suppose that  $\mathcal{A}(F)$  is not linear homogeneous, and prove that less than an  $|\mathcal{F}|^{-1/2}$  fraction of the equations are satisfied. Define for every  $y \in \mathcal{F}^{d(F)}$  the [r(F), 1]-LDF  $\phi_y$  as in the proof of Claim 5.12:

$$\forall t \in \mathcal{F} \qquad \phi_y(t) \doteq \mathcal{A}(F[ty])$$

As proven there,  $\phi_y$  is linear-homogeneous for less than an  $(|\mathcal{F}|^{-1/2})/2$  fraction of the points y.

Consider a point y for which  $\phi_y$  is not linear-homogeneous. Since  $t\mathcal{A}(F[y])$  is linear-homogeneous as a function of t,  $t\mathcal{A}(F[y]) \neq \mathcal{A}(F[ty])$  for all but at most an  $\frac{r(F)}{|\mathcal{F}|} \leq (|\mathcal{F}|^{-1/2})/2$  fraction of the t's, and hence at most an  $(|\mathcal{F}|^{-1/2})/2$  fraction of the equations  $\chi[y, t]$  are satisfied.

We have shown that for at least a  $1 - (|\mathcal{F}|^{-1/2})/2$  fraction of the y's  $\phi_y$  is not linear-homogeneous, and that for such y's  $\chi[y,t]$  is satisfied for less than an  $(|\mathcal{F}|^{-1/2})/2$  fraction of the t's. It follows that less than an  $|\mathcal{F}|^{-1/2}$  fraction of the equations are satisfied, as we needed to show.

<sup>¶</sup> f is homogeneous of degree m iff  $f(ty) = t^m f(y)$  for every y and t.

**The curve**  $\Gamma$ **.** Write the active part of  $\psi$  as

$$\psi_{\star} : \sum_{i=1}^{D} \alpha_i E[x_i] \tag{9}$$

As in the curve-extension representation-procedure, we define a curve  $\Gamma : \mathcal{F} \to \mathcal{F}^d$ which goes through the points associated with the active variables of  $\psi$ .

**Definition 18** (the curve of  $\psi$ ). Let  $\mathcal{H}_{sD-1} = \{a_1, \ldots, a_{sD}\}$  be an arbitrary subset of  $\mathcal{F}$ . Define  $\Gamma : \mathcal{F} \to \mathcal{F}^d$  to be the vector of (D-1)-degree polynomial functions satisfying

$$\forall \ 1 \le i \le D \qquad \Gamma(a_i) = x_i$$

Given an assignment  $\mathcal{A}$  for E, the assignment of F is used as an encoding of  $\mathcal{A}(E) \circ \Gamma$ . Unlike in the curve-extension representation-procedure, the correct encoding here is a linear-homogeneous LDF.

**The encoding.** The procedure generates a curve-verifier, whose equations are only satisfied if the assignment of F is the correct encoding of  $\mathcal{A}(E) \circ \Gamma$ . To define what the correct encoding is, suppose E is assigned an *s*-degree LDF g. The LDF  $g \circ \Gamma : \mathcal{F} \to \mathcal{F}$ , which is to be encoded by the assignment of F, has degree at most sD - 1. Its encoding is the following linear-homogeneous LDF,  $L_g$ :

$$\forall (y_1, \dots, y_{sD}) \in \mathcal{F}^{sD} \qquad L_g(y_1, \dots, y_{sD}) \doteq \sum_{i=1}^{sD} y_i g(\Gamma(a_i)) \tag{10}$$

The next claim shows how  $g \circ \Gamma$  can be reconstructed, given  $L_q$ .

**Claim 5.14** (linearizing-interpolation). Let g be an [s,d]-LDF, and for  $i = 1, \ldots, sD$ , let  $\gamma_i$  be the [sD - 1, 1]-LDF satisfying  $\gamma_i(a_i) = 1$  and  $\gamma_i(a_j) = 0$  for every  $j \neq i$ .

Then the polynomial vector function  $\widehat{\gamma} = (\gamma_1, \ldots, \gamma_{sD})$  satisfies  $L_g \circ \widehat{\gamma} = g \circ \Gamma$ , where  $L_g$  is as defined in Equation 10.

*Proof.*  $L_g$  is linear, hence  $L_g \circ \widehat{\gamma}$  is of degree at most sD - 1. Since it follows from the definition of  $L_g$  that  $L_g \circ \widehat{\gamma}(a_i) = g(\Gamma(a_i))$  for  $i = 1, \ldots, sD$ , we obtain that  $L_g \circ \widehat{\gamma} = g \circ \Gamma$  (also recall that  $g \circ \Gamma$  is of degree at most sD - 1).

**Generating the curve-verifier.** It follows from Claim 5.14 that if an assignment  $\mathcal{A}$  assigns a good LDF to E and assigns its encoding to F, then  $\mathcal{A}(F[\widehat{\gamma}(x)]) = \mathcal{A}(E[\Gamma(x)])$  for every  $x \in \mathcal{F}$ . To verify that F is assigned a correct encoding, the representation-procedure generates one equation  $\chi[x]$  in the curve-verifier for every  $x \in \mathcal{F}$  as follows:

$$\chi[x]$$
 :  $F[\widehat{\gamma}(x)] = E[\Gamma(x)]$ 

where the vector-function  $\hat{\gamma}$  is as defined in Claim 5.14.

The following claim shows that indeed the curve-verifier equations are not satisfied unless the assignment for F is a correct encoding, in the sense that  $\mathcal{A}(F) \circ \widehat{\gamma} = \mathcal{A}(E) \circ \Gamma$ . It is assumed that F is assigned a linear LDF, since otherwise the linearization-verifier equations cannot be satisfied.

**Claim 5.15.** Let  $\mathcal{A}$  be a feasible assignment for E and F, assigning a linear homogeneous LDF to F. Then less than an  $|\mathcal{F}|^{-1/2}$  fraction of the curve-verifier equations are satisfied unless  $\mathcal{A}(F) \circ \widehat{\gamma} = \mathcal{A}(E) \circ \Gamma$ . In the latter case all of the curve-verifier equations are satisfied, and moreover,

$$\forall (y_1, \dots, y_{sD}) \in \mathcal{F}^{sD} \qquad \mathcal{A}(F[y_1, \dots, y_{sD}]) \doteq \sum_{i=1}^{sD} y_i \mathcal{A}(E[\Gamma(a_i)]) \tag{11}$$

Proof. Assume that at least an  $|\mathcal{F}|^{-1/2}$  fraction of the equations  $\chi[x]$  are satisfied. This means that  $\mathcal{A}(F[\widehat{\gamma}(x)]) = \mathcal{A}(E[\Gamma(x)])$  for at least an  $|\mathcal{F}|^{-1/2}$  fraction of the x's. Since  $\mathcal{A}(F) \circ \widehat{\gamma}$  is an LDF of degree at most  $sD - 1 < |\mathcal{F}|^{-1/2}$  and  $\mathcal{A}(E) \circ \Gamma$  is an LDF of degree at most  $r(D-1) < |\mathcal{F}|^{-1/2}$ , this implies that  $\mathcal{A}(F) \circ \widehat{\gamma} = \mathcal{A}(E) \circ \Gamma$ .

In this case it follows that  $\mathcal{A}(F[\widehat{\gamma}(x)]) = \mathcal{A}(E[\Gamma(x)])$  for every  $x \in \mathcal{F}$ , and hence the equation  $\chi[x]$  is satisfied. In addition, since in particular  $\mathcal{A}(F[\widehat{\gamma}(a_i)]) = \mathcal{A}(E[\Gamma(a_i)])$  for every  $a_i, i = 1, \ldots, sD$ , one obtains from the definitions of  $\Gamma$  and  $\gamma$  that Equation 11 holds for all unit vectors. By the linearity of the assignment for F, it follows that Equation 11 holds for all points.

**Generating**  $\psi'$ . The procedure now generates the equation  $\psi'$  from  $\psi$  by replacing its active part by just one variable. Specifically,  $\psi'$  is obtained from  $\psi$  by removing  $\psi_{\star}$  and replacing it by  $F[y_*]$ , where  $y_* \doteq (\alpha_1, \alpha_2, \ldots, \alpha_D, 0, 0, \ldots, 0)$ ,

and the  $\alpha_i$ 's are the coefficients that appear in the active part of  $\psi$  (see Equation 9). The rational behind this replacement is explained by the following immediate claim.

**Claim 5.16.** Let  $\mathcal{A}$  be an assignment for E and F which satisfies Equation 11 in Claim 5.15. Then the value of  $\mathcal{A}(F[y_*])$  is the same as the evaluation of  $\psi_{\star}$ .

**Generating**  $\mathcal{E}_{\psi}$ . The linearization representation-procedure constructs the set of conjunctions  $\mathcal{E}_{\psi}$  as follows. For each triplet (x, y, t) where  $x, t \in \mathcal{F}$  and  $y \in \mathcal{F}^{d(F)}$ ,  $\mathcal{E}_{\psi}$  will have the conjunction of  $\psi'$ , the curve-verifier equation  $\chi[x]$ , and the linearization-verifier equation  $\chi[y, t]$ .

### Correctness of the algorithm.

The domain F that is generated by the linearization representation-procedure has the required parameters, and it is easy to verify that the running time is polynomial in |F|. To complete the proof of Lemma 5.5 let us show that  $\mathcal{E}_{\psi}$ has the extension and restriction properties, as the other required properties are obvious.

• Extension: Let  $\mathcal{A}$  be a good assignment for the variables of  $\Psi$ , that satisfies  $\psi$ . Let g be the [s, d]-LDF assigned to E, and extend  $\mathcal{A}$  to F by assigning  $L_g$  to it. We need to show that the extended  $\mathcal{A}$  satisfies the conjunctions of  $\mathcal{E}_{\psi}$ . According to the construction, it is enough to show that  $\psi'$  is satisfied and that the curve-verifier and linearization-verifier equations are satisfied as well.

Since  $\mathcal{A}(F) = L_g$  is a linear-homogeneous LDF, the linearizationverifier equations are satisfied by Claim 5.13. Also, g is an s-degree LDF, so by Claim 5.14

$$\mathcal{A}(F) \circ \widehat{\gamma} = L_q \circ \widehat{\gamma} = g \circ \Gamma = \mathcal{A}(E) \circ \Gamma$$

From Claim 5.15 we thus have that all of the curve-verifier equations are satisfied. Moreover, Claim 5.15 also implies that Equation 11 is satisfied, from which, by Claim 5.16, it follows that the evaluations of  $\psi_{\star}$  and of  $F[y_*]$  are equal. Since  $\psi$  is satisfied, and  $\psi'$  differs from  $\psi$ only in the substitution of  $\psi_{\star}$  by  $F[y_*]$ , we obtain that  $\psi'$  is satisfied as well. • Restriction: Let  $\mathcal{A}$  be a feasible assignment for the variables of  $\Psi$  and for F. We assume that these assignments satisfy at least an  $|\mathcal{F}|^{-1/2}$  fraction of the conjunctions in  $\mathcal{E}_{\psi}$ . This implies that  $\psi'$  is satisfied, and that at least an  $|\mathcal{F}|^{-1/2}$  fraction of the curve-verifier equations are satisfied, as well as an  $|\mathcal{F}|^{-1/2}$  fraction of the linearization-verifier equations. Let us prove that  $\psi$  is satisfied.

Since at least an  $|\mathcal{F}|^{-1/2}$  fraction of the linearization-verifier are satisfied, we gather from Claim 5.13 that F is assigned a linear-homogeneous LDF. Now Claim 5.15 implies that Equation 11 holds, and therefore by Claim 5.16,  $\psi_{\star}$  has the same evaluation as  $F[y_*]$ . Since  $\psi'$  is satisfied, this implies that  $\psi$  is satisfied as well.

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