Leaders

Autografting as first line treatment for chronic myeloid leukaemia

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Abstract

Interest in autografting for chronic myeloid leukaemia and its clinical relevance has revived in recent years. This followed observations that with various chemotherapeutic regimens it was possible to achieve, temporarily at least, peripheral blood and bone marrow that were Philadelphia negative. Bone marrow or peripheral blood progenitor cells could then be harvested and reinfused following a high dose procedure, hopefully eliminating any residual disease, and resulting in prolonged disease free survival. This ideal has not yet been successfully achieved with current strategies. Recent results indicate that eliminating residual disease with current chemotherapy is not normally achievable. The use of more sensitive technologies such as polymerase chain reaction has revealed persistent disease in most if not all apparently Philadelphia negative cases. This is confirmed by results where disease relapse occurs following transplant in these cases. Despite this, clinically relevant remissions are obtained and further trials are indicated. In this review present treatment is discussed and future strategies, using novel techniques as an adjunct to current treatment, are proposed that might improve on present results or even lead to the elusive goal of cure.

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The initial management of patients who present with chronic myeloid leukaemia (CML) is one of the more complex problems in the treatment of the haematological malignancies. For many years the mainstay of treatment was busulphan, an alkylating agent administered orally, which has gradually been replaced over the past decade by hydroxyurea. During this time α interferon has been shown to prolong the duration of the chronic phase, which neither hydroxyurea nor busulphan have been shown to do. A recent study from a French collaborative group suggests that the addition of subcutaneous cytarabine to α interferon for 10 days each month for six or nine months may prolong survival further. In the 1980s it was shown that allogeneic transplantation from a histocompatible sibling donor could cure a proportion of patients with CML. If a sibling is available, transplantation is considered the treatment of choice and is performed optimally in the first 12 months after diagnosis. Unfortunately, only a minority of patients may benefit from a sibling allogeneic transplant as a donor is available for only one in three patients, and elderly patients or those with intercurrent illnesses may be considered unsuitable for the procedure.

Until relatively recently the use of HLA matched unrelated donors was reserved for the youngest and fittest patients because of the daunting morbidity and mortality of such transplants. However, since the advent of molecular typing for both class I and class II major histocompatibility loci it is possible to select more precisely compatible donors. Evidence is beginning to accrue that patients who receive a bone marrow transplant from an unrelated donor who is fully matched at the molecular level may have the same outcome as if the donor was a histocompatible matched sibling. However, a proportion of patients will still not have a suitable donor or will be considered unfit to undergo the rigours of an allograft.

The role of autografting

The concept of using the patient’s own haematopoietic stem cells, in the form of an autologous marrow transplant, was developed in the 1970s. Spiers and colleagues at the Hammersmith Hospital in London, UK developed the concept of “perpetual reinduction of chronic phase”. They envisaged a situation where chronic phase stem cells would be collected from the blood at diagnosis and stored in liquid nitrogen. At the time of disease progression chemotherapy would induce remission and the patient would have chronic phase cells reinfused. Although this is an attractive notion in theory, resistance of the disease to chemotherapeutic agents has prevented this strategy from being successful. During the 1980s, despite sibling donor transplants being the treatment of choice, some exploratory studies were undertaken using autografting for CML while
the patient was still in chronic phase. These studies were begun at a time when unrelated donor panels were in their infancy and the risks of unrelated donor transplants were daunting. Autografting seemed to offer a low risk alternative to unrelated donor bone marrow transplantation, even if the likelihood of success was unclear.

Results from Goldman’s group at the Hammersmith Hospital were encouraging.25 They treated 14 patients who were in chronic phase of six to 44 months duration. Eleven elective autologous bone marrow transplants were done and the other three patients had been rescued with autologous cells after rejection of an allogeneic transplant. The procedure was well tolerated with two deaths, one not related to the autograft. At the time of the report 12 of the 14 patients were alive at a median follow up of 41 months (range 24–53) after transplantation; however, all but two patients were entirely Philadelphia positive.

In 1994 McGlave et al combined data from eight transplant groups who had been autografting in various guises from 1984 to 1992.11 Criteria for inclusion in the study was that they were consecutive CML autografts in each centre. Follow up ranged from 1–91 months (median 30). The data are somewhat confounded by the wide range of protocols used but the overall results showed, perhaps surprisingly, that of the 142 patients autografted in first chronic phase, median survival had not been reached at >30 months with 58% of patients still alive, the longest survivor at 91 months. This meta-analysis also provided some support for autologous transplants in accelerated phase patients in whom median survival was 36 months, but it did not provide any support for autotransplant in blast crisis patients who had a median survival of only 4.1 months.11

Experience gleaned over the past 10 years suggests two essential requirements before a patient with CML may be cured. First, they must receive cytoreductive treatment, which usually involves some form of transplant conditioning with busulphan, cyclophosphamide or cyclophosphamide and total body irradiation. A few patients may become Philadelphia negative after treatment with α interferon.25 Second, there should be some degree of graft versus leukaemia. Graft versus leukaemia was first identified in CML when a very high relapse rate was observed in patients undergoing allogeneic transplants from sibling donors in which the T lymphocytes had been removed from the transplant—the relative risk of relapse was 5.4 times (p value of <0.0001).12 T cell depletion is an effective way of controlling or preventing graft versus host disease following allogeneic bone marrow transplant, but in CML these benefits are offset by the increased relapse risk. Among patients who had received a T cell depleted bone marrow transplant results showed that those who had no acute graft versus host disease at all had an especially high rate of relapse.13 Importance of T lymphocytes for the maintenance of remission following transplantation for CML was confirmed when it proved possible to reinude remission by infusing lymphocytes from the original donor. It is also possible to achieve a similar effect by sudden cessation of immunosuppressive treatment, thereby releasing immune competent cells into the circulation.15 16

In the 1990s Carella and colleagues in Genoa, Italy, gave autografting in CML a new lease on life when they found that patients with CML regenerated with wholly, or more usually partially, Philadelphia negative haemopoiesis following intensive chemotherapy.17 18 This followed the observation that there was evidence of residual Philadelphia negative cells in many patients with CML, at least in the early stages of their disease.19 20 Although the Philadelphia negativity was generally not sustained, Carella’s group showed that it was possible, in the early recovery phase, to collect sufficient Philadelphia negative haemopoietic progenitor cells from peripheral blood to perform a subsequent autologous transplant (fig 1).

They used a highly intensive chemotherapy regimen of idarubicin 8 mg/m²/day iv push on days 1–5, cytarabine 800 mg/m² two hour infusion daily on days 1–5, and etoposide 150 mg/m² two hour infusion daily on days 1–3.21 The haemopoietic growth factor granulocyte colony stimulating factor was used to stimulate recovery of the neutrophils before peripheral blood stem cell collection. Carella’s initial series involved accelerated phase or blast crisis patients and although a significant proportion converted to Philadelphia negativity this was transient and all patients eventually relapsed. They subsequently moved on to explore the possibility of using this approach in 23 patients who were in chronic phase. Twenty of those were Philadelphia negative in blood after mobilisation and they engrafted well; seven remained Philadelphia negative two to 45 months after transplantation. Within this group of patients only three have died, none directly related to the transplant.22

Other groups took up the challenge to reproduce these findings. Our own group used a modified version of the Carella protocol in which only idarubicin and cytarabine were given to the patient. It was possible to collect peripheral blood stem cells from 23 of 41 patients from five centres who underwent this regimen that had either a complete or a major (<35% Philadelphia positive) cytogenetic response. A proportion of patients were also negative by Southern blotting but all patients were positive using polymerase chain reaction for bcr-abl. This study showed that there was

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**Figure 1** Approach adopted by Carella for autografting in CML. Initially mobilising late in disease, progression with ICE, but better results obtained by autografting in earlier chronic phase.
A potential therapeutic option involves harvesting cells in first chronic phase (CP1) and obtaining disease control by conventional chemotherapeutic means. This treatment would be augmented by vaccinating against bcr-abl thereby inducing an immune response against the malignant clone that could be further augmented by return of in vitro manipulated lymphocytes directed against bcr-abl.

One problem with peripheral stem cells collected in this way is that their quality is often poor—that is, the number of colony forming units and CD34 positive cells is often relatively few. Of 28 patients in our series who underwent autologous peripheral blood stem cell transplants three died within six months of the procedure. Nineteen remain alive with three in accelerated phase, three with quiescent disease on no treatment, and the rest alive in first chronic phase receiving hydroxyurea with or without α interferon. Median survival following transplant is still only 24 months although median survival from diagnosis in this highly selected series of patients is 55 months.

When to autograft?

It is clear that there is a case to be made for autografting as early treatment in chronic myeloid leukaemia. Most people would prefer to control the patient's white count with a drug such as hydroxyurea before proceeding to intensive treatment, rather than give high dose chemotherapy straight away. What is clear is that autografting in CML remains an experimental procedure the benefits of which are unknown. It is therefore important that any patients who receive such treatment should be within a trial or study in which the outcome is capable of being assessed. Ideally this would be in a randomised controlled trial such as the study being undertaken by the UK Medical Research Council in its CML IV trial. This trial is comparing a course of intensive treatment (idarubicin and cytarabine) followed by a bone marrow transplant following recovery from the intensive treatment with continuing interferon. The doses of chemotherapy, the transplant conditioning, and cells to rescue the patients from that conditioning have all been carefully chosen to minimise mortality and reduce morbidity. Suitable patients are those with no compatible sibling, or those too old for a sibling allograft and yet fit enough for intensive treatment. In our institution this would include all patients over the age of 60 who are fit and some patients between the ages of 50 and 60. It would be inappropriate to advise patients that they could have an autograft, with intensive conditioning such as high dose busulfan, and then at some later date be considered fit for unrelated donor transplant using similar or more intensive conditioning. It is important to give patients complete advice and make a decision as to the appropriate management at the beginning. Within the MRC trial patients receive their chemotherapy followed by the autograft starting three months from diagnosis after an initial trial of α interferon. Other national and international trials are investigating earlier autografting and/or the use of peripheral blood stem cell autografts.

There are few data on the optimal drugs and dosage regimens for the initial priming chemotherapy. We also do not know which growth factor or growth factor combinations are the most appropriate for mobilisation.

Other important issues to be considered in autologous transplantation relate to purging—a technique by which overt or potential malignant cells are removed from the transplant inoculum. At present such procedures are highly experimental and none has so far been proved effective. Deisseroth’s group has shown, using gene marking studies, that malignant cells in the CML rescue marrow do contribute to relapse although this is not the same as saying that they cause the relapse.

The future

It seems clear that increasingly intensive treatment for CML is not the whole answer. Despite sterling work by many groups around the world we will still benefit from improved understanding of the Philadelphia positive progenitor cells, in terms of biology and susceptibilities to specific means of disruption. Ideally autografts should stimulate the graft versus leukaemia effect. Techniques under consideration include dendritic cell or T lymphocyte infusions, prior vaccination against bcr-abl protein products, and, in the more distant future, DNA or RNA vaccines. Some pilot studies of antisense gene treatment targeted at blocking the bcr-abl locus have also been attempted although reproducibility remains a problem. Other target genes such as c-myb are also being investigated. Improved vectors and delivery systems appear likely to be required for gene therapy before antisense is likely to have a role in CML.

One view of the future would be to modify the schema proposed over 20 years ago by Spiers. Patients would, following diagnosis, have stem cells collected from which dendritic cells or specific T cell subsets would be isolated and amplified. Such cells together with unmanipulated stem cells could then be frozen for future use. Initial disease control would be
Autografting as first line treatment for chronic myeloid leukaemia

95

achieved following which some form of cyto-
duction (possibly with intensive treatment or α interferon) would be done until the patient had minimal residual disease—that is, disease that could not be detected conventionally. They may previously have been vaccinated against bcr-abl and could again be vaccinated once the disease was in a form of remission. Further maintenance of sustained disease free survival would be achieved by repeated infusions of dendritic cell and T cell combinations primed to seek out and destroy those cells with the bcr-abl gene expression. Stem cells to re-establish haemopoiesis may indeed be unnec-
sary (fig 2).

Allografting remains the treatment of choice when practical. If an autologous donor is unavailable autografts are currently a viable alternative and by preference should be performed early. This appears to induce a prolonged chronic phase, but the role of autografting in achieving a “cure” may be by returning selected and manipulated host cells capable of re-establishing a Philadelphia negative haemopoiesis and eliminating any Phila-
delphia positive clones with a graft versus leu-
kaemia effect.

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