A REVIEW ON ETIOPATHOGENESIS AND DRUGS USED FOR THE TREATMENT OF LEUKEMIA

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Abstract

Leukemia encompasses a clinically and pathologically diverse set of conditions whose incidence and prevalence are rising. In the past, leukemia was classified on the basis of the morphological characteristics of abnormally proliferating leucocytes in the blood and bone marrow. The etiology of leukemia mainly observed as Abnormalities of chromosomes have been found in various types of leukemia & other factors are like environmental factors, irradiation, and certain chemicals. This is a two part review with the first part focusing on the etiopathogenesis. The second part focused on management of Leukemia with drugs. For the management of leukemia the available treatments are chemotherapy, radiation therapy, targeted drug therapy, biological therapy and stem cell transplantation.

Keyword: Leukemia, Chemotherapy, Leucocytes, Stem cell Transplantation

1. INTRODUCTION

1.1. Leukemia

Leukemia is malignancy of the body’s blood-framing tissues, including the bone marrow and the lymphatic framework. There are numerous sorts of leukemia exist. A few types of leukemia are most normal in kids. Different types of leukemia happen primarily in grown-ups. Leukemia more often than not includes the white platelets. Your white platelets are powerful contamination contenders — they typically develop and partition in a methodical manner, as your body needs them. In any case, in the general population with leukemia, the bone marrow produces unusual white platelets, which don’t work appropriately. Treatment for leukemia can be mind boggling — relying upon the kind of leukemia and different elements. Yet, there are techniques and assets that can make your treatment fruitful.

1.2. Symptoms of Leukemia

Leukemia side effects change, contingent upon the kind of leukemia. Normal leukemia signs and side effects include:

➢ Fever or chills
➢ Persistent weariness, shortcoming
➢ Frequent or extreme diseases
➢ Losing weight easily
➢ Swollen lymph hubs, expanded liver or spleen
➢ Easy draining or wounding
➢ Recurrent nosebleeds
➢ Tiny red spots in your skin (petechiae)
➢ Excessive perspiring, particularly during the evening
➢ Bone agony or delicacy

2. ETIOLOGY

Researchers don’t comprehend the careful reasons for leukemia. It appears to create from a mix of hereditary and ecological components. How leukemia shapes as a rule, leukemia is thought to happen when some platelets get transformations in their DNA — the guidelines inside every cell that manage its activity. There might be different changes in the cells that presently can’t seem to be completely comprehended could add to leukemia. Certain anomalies cause the cell to develop and isolate all the more quickly and to keep
living when ordinary cells would bite the dust. After some time, these irregular cells can swarm out solid platelets in the bone marrow, prompting less sound white platelets, red platelets and platelets, causing the signs and side effects of leukemia [1, 2, and 3].

3. PATHOPHYSIOLOGY

The hematopoietic undifferentiated organism is in charge of delivering many platelets in the human body. The procedure begins when juvenile impact cells, lymphoid or myeloid, are discharged with huge cores that in the end become full grown and concentrated platelets with compacted DNA and littler cores. In leukemia, the youthful impact cell can’t advance in development due to various or single quality transformations that happen in the DNA. Different quality changes are the consequence of chromosome translocation, which is a mistake amid the cell division process. Single quality changes emerge from presentation to cancer-causing agents or radiation that causes a detached transformation. On account of transformations, the leukemia impact cells wildly recreate in the bone marrow and dominate. This is an issue, in light of the fact that the ordinary impact cells can’t utilize assets in light of constrained space. In this way, they are unfit to make adult specific cells, for example, platelets, red platelets, and white platelets. In constant leukemia, incompletely full grown cells create over an extensive stretch and don’t function as adequately. The most widely recognized causes are chromosomal anomalies, for example, cancellations, translocation, and trisomy. CML is an aftereffect of cells separating too rapidly, while CLL is a consequence of faulty apoptosis [4, 5].

4. LEUKEMIA CLASSIFICATION

4.1. Acute leukemia

In intense leukemia, the anomalous platelets are juvenile platelets (impacts). They can’t do their ordinary capacities, and they increase quickly, so the infection exacerbates rapidly. Intense leukemia requires forceful, convenient treatment.

4.2. Chronic leukemia

There are numerous sorts of unending leukemia’s. Some produce such a large number of cells and some reason too couple of cells to be created. Perpetual leukemia includes progressively adult platelets. These platelets reproduce or amass all the more gradually and can work regularly for a timeframe. A few types of interminable leukemia at first produce no early manifestations and can go unnoticed or undiscovered for quite a long time. The second kind of grouping is by sort of white platelet influenced:

4.3. Lymphocytic leukemia

This kind of leukemia influences the lymphoid cells (lymphocytes), which structure lymphoid or lymphatic tissue. Lymphatic tissue makes up your insusceptible framework.

4.4. Myelogenous leukemia

This kind of leukemia influences the myeloid cells. Myeloid cells offer ascent to red platelets, white platelets and platelet-creating cells.

4.5. Types of leukemia

The real kinds of leukemia are:

4.5.1. Acute lymphocytic leukemia (ALL)

This is the most well-known sort of leukemia in youthful youngsters. ALL can likewise happen in grown-ups.

4.5.2. Acute myelogenous leukemia (AML)

AML is a typical sort of leukemia. It happens in kids and grown-ups. AML is the most widely recognized kind of intense leukemia in grown-ups.

4.5.3. Chronic lymphocytic leukemia (CLL)

With CLL, the most widely recognized incessant grown-up leukemia, you may feel well for a considerable length of time without requiring treatment.

4.5.4. Chronic myelogenous leukemia (CML)

This kind of leukemia basically influences grown-ups. An individual with CML may have few or no side effects for
a considerable length of time or years before entering a stage in which the leukemia cells develop all the more rapidly. Other, rarer kinds of leukemia exist, including bristly cell leukemia, myelodysplastic disorders and myeloproliferative issue.

5. RISK FACTORS

Variables that may expand your danger of building up certain sorts of leukemia include:

1) Previous disease treatment. Individuals who’ve had particular sorts of chemotherapy and radiation treatment for different diseases have an expanded danger of building up specific kinds of leukemia.

2) Genetic disarranges. Hereditary anomalies appear to assume a job in the advancement of leukemia. Certain hereditary issue, for example, Down disorder, are related with an expanded danger of leukemia.

3) Exposure to specific synthetic compounds. Presentation to specific synthetic substances, for example, benzene — which is found in gas and is utilized by the concoction business — likewise is connected to an expanded danger of certain sorts of leukemia.

4) Smoking. Smoking cigarettes builds the danger of intense myelogenous leukemia.

5) Family history of leukemia. On the off chance that individuals from your family have been determined to have leukemia, your hazard for the malady might be expanded.

6. TESTS AND DIAGNOSIS

Specialists may discover constant leukemia in a standard blood test, before side effects start. On the off chance that this occurs, or on the off chance that you have signs or side effects that recommend leukemia, you may experience the accompanying indicative tests:

1) Physical test. Your specialist will search for physical indications of leukemia, for example, fair skin from paleness, swelling of your lymph hubs, and development of your liver and spleen.

2) Blood tests. By taking a gander at an example of your blood, your specialist can decide whether you have irregular dimensions of white platelets or platelets — which may recommend leukemia.

3) Bone marrow test. Your specialist may prescribe a method to expel an example of bone marrow from your hipbone. The bone marrow is evacuated utilizing a long, meager needle. The example is sent to a research center to search for leukemia cells. Particular trial of your leukemia cells may uncover certain attributes that are utilized to decide your treatment alternatives.

4) You may experience extra tests to affirm the determination and to decide the kind of leukemia and its degree in your body. Particular sorts of leukemia are arranged into stages, showing the seriousness of the infection. Your leukemia’s stage enables your specialist to decide a treatment plan.

7. TREATMENTS AND DRUGS

Treatment for your leukemia depends on many factors. Your doctor determines your leukemia treatment options based on your age and overall health, the type of leukemia you have, and whether it has spread to other parts of your body. Common treatments used to fight leukemia include:

7.1. Chemotherapy

Chemotherapy is the significant type of treatment for leukemia. This medication treatment utilizes synthetic substances to execute leukemia cells. Contingent upon the kind of leukemia you have, you may get a solitary medication or a mix of medications. These medications may arrive in a pill structure, or they might be infused straightforwardly into a vein.

7.2. Biological therapy

Organic treatment works by utilizing medicines that help your invulnerable framework perceive and assault leukemia cells.

7.3. Targeted therapy

Directed treatment utilizes drugs that assault explicit vulnerabilities inside your malignancy cells. For instance, the medication imatinib (Gleevec) stops the activity of a
protein inside the leukemia cells of individuals with unending myelogenous leukemia. This can help control the malady.

7.4. Radiation therapy

Radiation treatment utilizes X-beams or other high-vitality bars to harm leukemia cells and stop their development. Amid radiation treatment, you lie on a table while an expansive machine moves around you, guiding the radiation to exact focuses on your body. You may get radiation in one explicit region of your body where there is a gathering of leukemia cells, or you may get radiation over your entire body. Radiation treatment might be utilized to plan for an immature microorganism transplant.

7.5. Stem cell transplant

An immature microorganism transplant is a strategy to supplant your unhealthy bone marrow with sound bone marrow. Before an undifferentiated organism transplant, you get high dosages of chemotherapy or radiation treatment to crush your infected bone marrow. At that point you get an implantation of blood-shaping immature microorganisms that help to revamp your bone marrow. You may get undifferentiated organisms from a benefactor, or at times you might almost certainly utilize your very own foundational microorganisms. An immature microorganism transplant is fundamentally the same as a bone marrow transplant.

8. TREATMENT RESPONSE TERMINOLOGY

8.1. Complete Remission

- Absence of leukemia cells in the bone marrow
- All signs and symptoms are resolved
- No more than 5% of blast cells in the bone marrow or in the blood stream
- Normal blood cell counts

8.2. Refractory

Small amount of leukemia cells remain in the blood marrow after intense therapy, can be determined by minimal residual disease testing

8.3. Relapse

Leukemia cells return in the blood marrow after complete remission has been reached.

9. TREATMENT OF ACUTE LEUKEMIA

At the start, escalated mix chemotherapy is given in the expectation of accomplishing a total reduction (CR). This underlying period of treatment is named Induction or Remission Induction Chemotherapy. A CR must be accomplished by virtual removal of the bone marrow, trailed by recuperation of ordinary haemopoiesis. On the off chance that two cycles of treatment neglect to actuate CR, an elective medication routine can be utilized. On the off chance that this is fruitless, it is impossible that CR will be accomplished. The consequent span of the principal abatement is firmly connected to survival. Reduction is characterized as the nonattendance of all clinical and minuscule indications of leukemia, under 5% shoot structures in the bone marrow and return of typical cellularity and haemopoietic components. Regardless of accomplishing CR, mysterious remaining ailment (additionally named insignificant lingering sickness or MRD) will persevere, and further escalated treatment is given trying to support the abatement. This Post-Remission Consolidation Therapy may involve chemotherapy or a mix of chemotherapy and bone marrow transplantation.

10. Acute lymphocytic leukemia (ALL)

1. The blend of vincristine, prednisolone, anthracyclines and asparaginase instigates CR in about 90% of kids with ALL and 80% of grown-ups, albeit tragically backslide is unquestionably increasingly basic in grown-ups.
2. Other dynamic medications in the treatment of ALL incorporate methotrexate, 6-mercaptopurine, cyclophosphamide and mitoxantrone.
3. Patients with ALL are at a high danger of creating CNS invasion. Cytotoxic medications infiltrate inadequately into the CNS which in this manner goes about as an asylum site for leukaemic cells. Therefore, all patients with ALL get CNS prophylaxis. Cranial light in addition to intrathecal methotrexate or high portion foundational methotrexate can be utilized.
4. Maintenance treatment is critical to continue a CR. It is typically milder than enlistment or solidification chemotherapy, yet is carried on for something like year and a half. Treatment as a rule comprises of week by week methotrexate and every day 6-mercaptopurine with discontinuous vincristine and prednisolone.

5. The treatment of backslide infection differs with the site of backslide. Detached CNS or testicular backslide might be effectively treated with radiation and reinduction treatment. Fix can in any case be accomplished for certain patients. Bone marrow backslide is considerably more hard to fix, particularly in the event that it happens early.

6. A little extent of pediatric patients and a bigger proportion of grown-up patients have the Philadelphia chromosome translocation inside their ALL impacts. Such patients have a moderately poor anticipation and hence require increasingly serious treatment. There is some proof that medicate mixes including imatinib may improve the reaction of these leukaemias to treatment.

11. DRUG TREATMENT

11.1. Acute lymphocytic leukemia

11.1.1. Induction (4 weeks)

VPLD Regimen: Vincristine, Prednisolone, L-Asparaginase, Daunorubicin
Intensification (1 week)

VDPECT Regimen: Vincristine, Daunorubicin, Prednisolone, Etoposide, Cytarabine, Thioguanine

11.1.2. CNS prophylaxis (3 weeks)

Cranial irradiation by Methotrexate

11.1.3. Maintenance therapy (2 years)

MPV Regimen: Methotrexate, 6-Mercaptopurine, Prednisolone, Vincristine

11.2. Acute Myeloblastic Leukemia

Same as in ALL the treatment as follows for the AML

11.2.1. Induction Chemotherapy

ADE Regimen: Cytosine arabinoside, Daunorubium, Etoposide

11.2.2. Consolidation chemotherapy

MACE Regimen: Amsacrine, Cytosine arabinoside, Etoposide

MidAC Regimen: Mitoxantrone, Cytosine arabinoside

11.3. Chronic leukemia

Chronic leukemia is an increase of abnormal white blood cells. It differs from acute leukemia, and is categorized as myelogenous or lymphocytic.

11.3.1. Chronic Myelocytic Leukaemia

The treatment of CML has been essentially palliative, producing modest increases in survival, but with the main aim of keeping patients asymptomatic by normalizing the WBC. Hydroxycarbamide was the most widely used drug in the management of CML in chronic phase. Treatment with hydroxycarbamide is initiated at a dose of 1.5–2 g/day and usually brings the WBC under control within 1–2 weeks. The dose can then be reduced to a maintenance dose of 0.5–2 g/day. Withdrawing or reducing the dose abruptly can cause a rebound increase in WBC. The side effects of hydroxycarbamide are generally mild but include rashes and gut disturbances.

1. Tyrosine Kinase Inhibitors as initial therapy
   i. Imatinib
   ii. Bosutinib
   iii. Ponatinib

2. If initial therapy fails
   i. Interferon Alpha, Hydroxy urea, Cytarbine
   ii. Busulfan

3. Leukapheresis (Removal of White Blood Cells)

4. Stem Cell Transplant

11.3.2. Chronic lymphocytic leukemia

Patients who might not be able to tolerate the side effects of strong chemotherapy (chemo) are often treated with chlorambucil with a monoclonal antibody like obinutuzumab. Other options include the targeted drug ibrutinib (Imbruvica) alone, and
rituximab alone or with a corticosteroid like prednisone. There are other options, too. In stronger and healthier patients, commonly used treatments include:
1. FCR: fludarabine, cyclophosphamide, and rituximab
2. Bendamustine (sometimes with a CD20 monoclonal antibody)
3. Ibrutinib
4. FR: fludarabine and rituximab
5. High-dose prednisone and rituximab
6. PCR: pentostatin, cyclophosphamide, and rituximab
7. Alemtuzumab with rituximab

Other drugs or combinations of drugs may also be used.

12. RADIATION OR SURGERY

On the off chance that the main issue is an expanded spleen or swollen lymph hubs in a single piece of the body, restricted treatment with low-portion radiation treatment might be utilized. Splenectomy (medical procedure to expel the spleen) is another alternative if the augmented spleen is causing side effects.

13. LEUKAPHERESIS

Now and again exceptionally high quantities of CLL cells in the blood cause issues with typical course. This is called leukostasis. Chemo may not bring down the quantity of cells until a couple of days after the principal portion, so before the chemo is given, a portion of the phones may should be expelled from the blood with a system called leukapheresis. This treatment brings down blood tallies immediately. The impact endures just for a brief span, however it might help until the chemo gets an opportunity to work. Leukapheresis is likewise once in a while utilized before chemo if there are extremely high quantities of leukemia cells (notwithstanding when they aren't influencing issues) to anticipate tumor lysis disorder.

14. STEM CELL TRANSPLANT

Some people who have very high-risk disease (based on prognostic factors) may be referred for possible stem cell transplant (SCT) early in treatment.

15. SUPPORTIVE CARE

Dealing with the difficulties related with treatment is basic and similarly as significant as the treatment. There are numerous potential difficulties related with chemotherapy, including bladder and skin changes, wounding, obstruction, looseness of the bowels, exhaustion, male pattern baldness, sickness, neuropathy, unjustifiable dying, and retching. The antagonistic impact experienced decides the requirement for medicines, which incorporates antiemetic, granulocyte-province animating component, and transfusions. Different operators, for example, corticosteroids and anti-toxins, might be important to help battle contaminations in a smothered resistant framework. Notwithstanding oncology, these patients perpetually require an included interdisciplinary group of caseworkers, clinical drug specialists, irresistible illness experts, and nutritionists.

REFERENCES