Adverse reactions to targeted and non-targeted chemotherapeutic drugs with emphasis on hypersensitivity responses and the invasive metastatic switch

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Abstract More than 100 drugs are used to treat the many different cancers. They can be divided into agents with relatively broad, non-targeted specificity and targeted drugs developed on the basis of a more refined understanding of individual cancers and directed at specific molecular targets on different cancer cells. Individual drugs in both groups have been classified on the basis of their mechanism of action in killing cancer cells. The targeted drugs include proteasome inhibitors, toxic chimeric proteins and signal transduction inhibitors such as tyrosine kinase (non-receptor and receptor), serine/threonine kinase, histone deacetylase and mammalian target of rapamycin inhibitors. Increasingly used targeted vascular (VEGF) and platelet-derived endothelial growth factor blockade can provoke a range of pathological consequences. Many of the non-targeted drugs are cytotoxic, suppressing haematopoiesis as well as provoking cutaneous eruptions and vascular, lung and liver injury. Cytotoxic side effects of the targeted drugs occur less often and usually with less severity, but they show their own unusual adverse effects including, for example, a lengthened QT interval, a characteristic papulopustular rash, nail disorders and a hand–foot skin reaction variant. The term hypersensitivity is widely used across a number of disciplines but not always with the same definition in mind, and the terminology needs to be standardised. This is particularly apparent in cancer chemotherapy where anti-neoplastic drug-induced thrombocytopenia, neutropenia, anaemia, vascular disorders, liver injury and lung disease as well as many dermatological manifestations sometimes have an immune basis. The most insidious of all adverse consequences of targeted therapies, however, are tumour adaptation, increased malignancy and the invasive metastatic switch seen with anti-angiogenic drugs that inhibit the VEGF-A pathway. Adverse reactions to 44 non-targeted and 33 targeted, frequently used, chemotherapeutic drugs are presented together with discussions of diagnosis, premedications, desensitizations and importance of understanding the mechanisms underlying the various drug-induced reactions. There is need for wide-ranging acceptance of what constitutes a hypersensitivity reaction and for allergists to be more involved in the diagnosis, treatment and prevention of chemotherapeutic drug-induced hypersensitivity reactions.

Keywords Chemotherapy-induced hypersensitivities · Cancer drug hypersensitivities · Cancer drug adverse effects · Hypersensitivities to chemotherapy drugs · Drug allergies · Drug hypersensitivities · Drugs and metastatic switch

1 Adverse reactions to drugs

Forty years ago, The World Health Organization [1] defined an adverse drug reaction (ADR) as ‘a response to a drug that is noxious and unintended and occurs at doses normally used in man for the prophylaxis, diagnosis or therapy of disease, or for modification of physiological function’. This often-quoted definition and some others subsequently advanced have not always been universally accepted without criticisms especially with regard to the term ‘noxious’, the narrow use of ‘drug’, queries on just how minor an adverse reaction can be and other alleged ambiguities. While universal agreement on a
definition is unlikely, that proposed by Edwards and Aronson [2] seems adequate: ‘An appreciably harmful or unpleasant reaction, resulting from an intervention related to the use of a medicinal product, which predicts hazard from future administration and warrants prevention or specific treatment, or alteration of the dosage regimen, or withdrawal of the product’. Meanwhile, the US Food and Drug Administration (FDA) defines an adverse event as ‘any undesirable experience associated with the use of a medical product in a patient’ and states that any such serious event should be reported to the FDA [3]. A patient event is regarded as ‘serious’ when the outcome is death, life-threatening, hospitalisation, disability or permanent damage, congenital anomaly/birth defect, required intervention to prevent permanent impairment or damage (devices) and other serious important medical events (for example, allergic bronchospasm, serious blood dyscrasias or seizures or convulsions that do not result in hospitalisation).

ADRs have been classified into seven different reaction types A to G with group A termed augmented pharmacologic effects; B, bizarre; C, chronic (or continuous effects); D, delayed effects; E, end-of-treatment (or withdrawal) effects; F, failure of therapy and G, genetic reactions [2, 4]. For our purposes, types A and B, distinguished primarily on the basis of dose-related and non-dose-related reactions, respectively, are the reactions of main concern. Type A reactions are said to account for ~80 % of ADRs; reactions are predictable, can be anticipated from the drug’s pharmacological actions and are dose-dependent and resolve when the dose is reduced or withdrawn. Type B, or unpredictable reactions, sometimes called idiosyncratic drug reactions, are generally unrelated to the drug’s pharmacological actions and are independent of dose and usually (but not always) resolve when treatment is stopped. Type B reactions can be divided into immune-mediated or true allergic responses and non-immune, non-allergic sensitivities (Table 1). The immune-based category is made up of the true hypersensitivities defined by Gell and Coombs: type I, IgE antibody-mediated reactions; type II, cytotoxic reactions; type III, immune-complex-mediated hypersensitivities and type IV, delayed, cell-mediated responses [5]. Three type B non-immune drug sensitivities are recognized: pseudoallergy, idiosyncratic reactions and type B intolerances. Examples of the four allergic and three non-allergic sensitivities of type B ADRs are summarized side by side in Table 1. Many drugs that cause ADRs are not limited to one type of reaction, and some, such as the penicillins and non-steroidal anti-inflammatory drugs (NSAIDs), may be involved in up to four (including all four types of hypersensitivities) or even more different adverse responses.

2 Hypersensitivity reactions to drugs

The term hypersensitivity is a much misused term even within its ‘home’ discipline of allergy/immunology. In 2001, the European Academy of Allergology and Clinical Immunology (EAACI) published a EAACI Position Statement entitled, in part, ‘A revised nomenclature for allergy’ [6]. Following this lead, the World Allergy Organization (WAO) set about promoting what was described as acceptable nomenclature for allergic diseases with the stated ultimate goal of improving communication in the field of allergy. In 2003, the WAO Nomenclature Review Committee defined ‘hypersensitivity’ as ‘objectively reproducible symptoms and signs initiated by exposure to a defined stimulus at a dose tolerated by normal persons’ [7]. Far from improving communication (and apart from the absence of any reference to the adverse nature of a reaction), this definition creates confusion in the mind of workers familiar with the accepted Gell and Coombs classification based on four types of hypersensitivities. It is also difficult for those reading and attempting to understand the existing allergy/immunology literature where ‘hypersensitivity’ is generally taken to mean adverse signs and symptoms initiated by an antigenic stimulus that is tolerated by a ‘normal’ person and which has an immune component or basis. Long-term usage of the terms immediate, delayed, cell-mediated and types I, II, II and IV hypersensitivity has become in-grained and synonymous with immune reactions. Reactions to some drugs show features of a hypersensitivity response even though no immune mechanism has yet been identified. Examples include some (not all) reactions to contrast media [8] and NSAIDs where, in the latter case, drug-induced redirection of mediator synthesis in the arachidonic acid cascade from the cyclooxygenase to the lipoxygenase pathway occurs [9]. In these cases, such reactions can be described as ‘sensitivities’ or ‘tolerances’ rather than ‘non-allergic hypersensitivity’ recommended by the WAO. In terms of the Gell and Coombs definitions and classifications [5], ‘non-allergic hypersensitivity’ is essentially a contradiction in terms.

2.1 Use of the term ‘hypersensitivity’ in relation to chemotherapeutic drugs

Across all medical disciplines, and in the community at large, it is well-known that drugs used to treat cancers cause a range of detrimental side effects ranging from mild (e.g. appetite suppression) to severe (e.g. bone marrow suppression) and even death. Unfortunately, and for a number of reasons, medical/scientific classification of these reactions is often deficient, and this is particularly apparent for many adverse reactions incorrectly described as ‘hypersensitivities’ [10] and for others where that description was not applied when it was clearly appropriate. In the area of side effects of chemotherapeutic drugs then, the picture is confusing and in need of standardisation in both definitions and the use of terms. The term ‘hypersensitivity’ remains widely used, for example in the oncology literature, clinical trial reports, pharmaceutical
Table 1  Classification of immune (hypersensitivity) and non-immune type B adverse drug reactions

| Immune (allergic; hypersensitivity) | Drug examples | Non-immune (non-allergic sensitivities) |
|-------------------------------------|--------------|----------------------------------------|
| Hypersensitivity type¹              |              |                                        |
| Type I² (IgE antibody-mediated)     | Penicillins; cephalosporins; neuromuscular blockers; quinolones; pyrazolones; ppi; l-asparaginase; mAbs | Pseudoallergy³ Most reactions to NSAIDs. Direct mast cell degranulation (neuromuscular blockers; opioids; contrast media; vancomycin) |
| Type II⁴ (antibody [IgG, IgM]-mediated cytotoxic reactions) | Penicillins; cephalosporins; quinine; quinidine; sulphonamides; thiouracil; gold salts; oxaliplatin; fludarabine; irinotecan; suramin; mAbs | Idiosyncratic reactions⁵ Halothane hepatitis; malignant hypothermia; drug-induced anaemia in Glu-6-PO₄ dehydrogenase deficient individuals (antimalarials, sulphonamides, dapsone) |
| Type III⁶ (immune complex-mediated via IgG, IgM) | Penicillins; cephalosporins; sulphonamides; tetracycline; NSAIDs; carbamazepine; allopurinol; mAbs; oxaliplatin | Intolerance Tinnitus induced by small doses aspirin. Some reactions to contrast media and NSAIDs⁷ |
| Type IV⁸ (delayed; T cell-mediated) | NSAIDs; penicillins; other antimicrobials; anticonvulsants (carbamazepine); local anaesthetics; sulphonamides; hydroxychloroquine; cytarabine; tamoxifen; imatinib; mAbs |                                        |

Type B are so-called ‘bizarre’ reactions. Uncommon relative to type A (‘augmented’, pharmacologic effects) reactions, unpredictable, rarely dose-dependent and unrelated to drug’s pharmacologic action. Reactions show high mortality relative to other categories of ADRs

ppi proton pump inhibitors, NSAIDs non-steroidal anti-inflammatory drugs
1 Defined and classified according to Gell and Coombs [5]
2 e.g. anaphylaxis, bronchospasm, cardiovascular collapse, urticaria, angioedema
3 May closely resemble true type I reactions—termed anaphylactoid.
4 e.g. drug-induced haemolytic anaemia, immune thrombocytopenia, immune form of agranulocytosis
5 May be unrelated or related to dose. Uncommon, unpredictable, unrelated to drug’s pharmacologic action
6 e.g. serum sickness, drug-induced vasculitis
7 Reactions that are clearly not hypersensitivity responses, pseudoallergic or idiosyncratic in nature
8 e.g. allergic contact dermatitis, psoriasis, maculopapular exanthema, acute generalized exanthematous pustulosis, drug reaction with eosinophilia and systemic symptoms, erythema multiforme, fixed drug eruption, Stevens–Johnson syndrome, toxic epidermal necrolysis

company drug descriptions and pharmacological studies but clearly it does not always mean the same thing—no common definition has been adopted within and between disciplines. The word is often used to cover reactions that clearly have no immune basis and which are otherwise not easy to classify [11]. On the other hand, since mechanisms for many drug reactions are not known, generally not investigated and appropriate tests are either not at hand or even developed, some true hypersensitivities remain unrecognised and therefore incorrectly classified.

3 Chemotherapeutic drugs and their classification on the basis of mechanism of action

The number of drugs used for the treatment of the many different cancers is in excess of 100 and growing [12, 13] and this is reflected in the numbers approved by the FDA, those awaiting approval, entries in the National Cancer Institute (NCI) drug dictionary and new drugs being assessed in clinical trials [13, 14]. Many of the compounds used in chemotherapy for many years have cytotoxic properties arising from a limited number of different mechanisms of action. These drugs represent a wide range of chemical structures and include, amongst other actions, alkylating agents, antimetabolites, cytoskeletal disruptors and drugs that directly affect DNA or protein synthesis [15]. These are all agents with relatively broad, rather than targeted and specific, modes of action. Alkylating agents like busulfan, nitrogen mustards and the platinum-based agent cisplatin that cross-link DNA; antimetabolite nucleoside analogues like gemcitabine; inhibitors of mitotic cell division by taxanes and Vinca alkaloids; drugs that intercalate with DNA like actinomycin D; pyrimidine analogues including 5-fluorouracil and folate antimetabolites such as pemetrexed are typical examples of non-targeted drugs.
that contrast with drugs developed from current strategies based on a more refined understanding of individual cancers and a research focus on selected molecular targets for specific types of cancers [15]. Signal transduction inhibitors which include tyrosine kinase and mammalian target of rapamycin (mTOR) and proteasome inhibitors are examples of the latter approach [16]. The literature contains a number of different classifications of chemotherapeutic agents based on chemical structure (e.g. alkyl sulphonates, anthracyclines, diterpines, organoplatinum agents), mechanisms of action, broad pharmacological categories (e.g. antibiotics, hormones), plant origin (taxanes, *Vinca* alkaloids) and where drugs act in the cell cycle (S, M, G1 or G2 phase). In many published classifications, these different categories are mixed together, for example, as divisions into alkylating agents, antibiotics, topoisomerase inhibitors, drugs derived from plants etc. (see, for example, [17]) rather than a division based on a single criterion such as mechanism of action or chemical class. Of course, any classification based on a single shared factor/property for all chemotherapeutic drugs is not straightforward. Many of the drugs show more than one mechanism of action, chemical structures even within one group can be quite diverse (e.g. alkylating agents) and some features (e.g. being plant-derived) are seen in only a few compounds. The classification selected here is based firstly on the division into non-targeted (Table 2) and targeted drugs (Table 3) and then on what is currently known of each drug’s mechanism(s) of action.

### 3.1 Non-targeted drugs

The anti-cancer action of traditional drugs used for chemotherapy depends on their capacities to destroy rapidly dividing uncontrolled cancer cells via their cytotoxic properties or by causing the cells to undergo apoptosis. However, these drugs also harm normal rapidly dividing cells such as those in the gastrointestinal tract, bone marrow and hair follicles causing well-known side effects like mucositis, stomatitis, myelosuppression and alopecia. This absence of targeted specificity contrasts with the current strategy of selectively killing a particular type of cancer cell. Some of the earliest anti-neoplastic drugs were alkylating agents that act by damaging DNA. A number of chemically different types of alkylating chemicals such as nitrogen mustards (e.g. chlorambucil), alkyl sulphonates (busulfan), nitrosoureas (carmustine), triazines (dacarbazine) and ethylenimines (altretamine) (Table 2) are still administered to cancer patients. Anti-metabolites made up essentially of pyrimidine analogues (e.g. 5-fluorouracil), nucleoside analogues (gemcitabine) and anti-folates (methotrexate) form a second large group of what are still the most widely used cytostatic/cytotoxic drugs in cancer therapy. A third group, the mitotic inhibitors containing the important taxanes docetaxel and paclitaxel, inhibit mitotic cell division by binding to microtubules while L-asparaginase is an important example of a small number of drugs that affect leukemic cells by disrupting protein synthesis. A chemically diverse variety of structures comprising many different drugs is used for chemotherapy by virtue of the direct drug effects on DNA (Table 2). This group includes inhibitors of DNA synthesis such as hydroxyurea, a number of structurally different compounds (organoplatinum agents, the antibiotic mitomycin C and pyrrolobenzodiazepines [132]) that cross-link DNA, topoisomerase I and II inhibitors, drugs such as the antibiotic mitomycin C that intercalate between DNA base pairs, inhibitors of DNA methylation (azacitidine), bleomycin which induces breaks in DNA and inhibitors of the poly-ADP ribose polymerase (PARP) family of enzymes that are critical in the maintenance of DNA integrity and repair [133]. PARP inhibitors appear to trap PARP1 and PARP2 enzymes at damaged DNA sites. These trapped enzyme–DNA complexes block DNA replication and are highly toxic to cells. In fact, it has been suggested that PARP inhibitors should be assessed on their potency to trap PARP as well as on their capacities to inhibit the enzymes. As shown in Table 2, some chemotherapeutic drugs demonstrate more than one anti-tumour activity. For example, the platinum drugs which cross-link DNA and are sometimes classified along with the alkylating agents and the anthracycline antibiotics daunorubicin (or daunomycin) and doxorubicin (adriamycin) show both topoisomerase II inhibitory activity and also intercalate between base pairs in the DNA helix thus preventing DNA replication and protein synthesis. Lenalidomide [134], listed in Table 2 in the ‘miscellaneous’ group, is related to the teratogenic drug thalidomide. It has a number of different activities suggesting its application to a range of haematological and solid tumours.

### 3.2 Targeted drugs

Specific and effective targeting of tumour cells without accompanying toxicity has always been desired in cancer therapy, and this has started to be realized with the introduction of monoclonal antibodies (mAbs) and signal transduction therapies. The former proteinaceous agents, unlike pharmacologically active ‘small’ molecules, utilize precise immunological targeting of cancer cells and are not considered here. The principle of signal transduction therapies is shown diagrammatically in Fig. 1. Signal transduction [16] involves the utilization of biochemically induced signals generated by a range of large and small molecules such as growth factors, neurotransmitters, hormones, cytokines, chemokines and ATP into a wide variety of cell responses like cell division, metabolic changes, gene expression and cell death. Signal transduction therapy then depends on identifying signalling proteins and their altered pathways. The original employment of this approach for cancer therapy goes back to the use of tamoxifen as an oestrogen receptor antagonist in treating some forms of breast cancer that are dependent on oestrogen for their growth [135]. Since this early breakthrough with
| Drug generic and trade names and classification on the basis of mechanism | Cancer indications | Mechanism(s) of action | Hypersensitivity/adverse reactions | Systemic | Cutaneous | References |
|---|---|---|---|---|---|---|
| **Alkylating agents** | | | | | | |
| Alkyl sulphonates | | | | | | |
| Busulfan (Myleran®, Busulfex®) | CML, may be given prior to bone marrow/stem cell transplantation | Alkylation producing guanine-adenine cross-links | Interstitial pneumonia; pulmonary fibrosis; thrombocytopenia; leukopenia; anaemia; veno-occlusive disease | Urticaria; pruritus; alopecia; erythema multiforme; xerosis | | [18, 19] |
| Nitrogen mustards | | | | | | |
| Chlorambucil (Leukeran®) | CLL; NHL; WM | Alkylation of guanine at N-7 and cross-linking of DNA | Fever; thrombocytopenia; anaemia; neutropenia; rash; interstitial pneumonitis; GI; hepatotoxicity | Urticaria; contact dermatitis; exanthema; rash; erythema multiforme; stomatitis; ten | | [19-21] |
| Cyclophosphamide (Cytoxan®, Endoxan®, Neosar®) | Lymphomas; brain; leukemia | Alkylation of guanine at N-7 and cross-linking of DNA | Anaphylaxis; bronchospasm; bms; GI haemorrhagic cystitis | Urticaria; angioedema; acne; erythema multiforme; pigm (skin, nails); stomatitis; vasculitis; sjg; ten | | [19, 21-24] |
| Ifosfamide (Ifex®) | Breast; brain; bone; cervical; lung; ovarian; lymphomas | Alkylation and cross-linking of DNA | Bronchospasm; myelosuppression; thrombocytopenia; encephalopathy; nephrotoxicity; GI | Urticaria; stomatitis; alopecia; erythema; hyper pigm | | [21, 25] |
| Mechlorethamine (Mustargen®) | CML; CLL; CTCL; HL; NHL; brain | Alkylation of guanine at N-7 and cross-linking of DNA | Anaphylaxis; lymphopenia; GI | | | [21, 25] |
| Melphalan (Alkeran®) | MM; ovarian | Alkylation of guanine at N-7 and cross-linking of DNA | Anaphylaxis; interstitial pneumonitis; pulmonary fibrosis; bronchospasm; thrombocytopenia; encephalopathy; nephrotoxicity; GI | | | [26] |
| Nitrosoureas | | | | | | |
| Carmustine (BiCNU®) | Brain; MM; lymphomas | Alkylates DNA and RNA and cross-links DNA | Thrombocytopenia; leukopenia; anaemia; pulmonary fibrosis infiltrates; GI | | | [19] |
| Streptozocin (Zanosar®) | Brain; HL; islet cells pancreas | Translocated into pancreas β-cells via GLUTZ. Damages DNA and activates poly ADP-ribosylation | Anaphylaxis; interstitial pneumonitis; pulmonary fibrosis; bms | | | [27, 28] |
| Triazines | | | | | | |
| Dacarbazine (DTIC-Dome®) | Malign melanoma; HL; sarcoma; islet cells pancreas; neuroblastoma | 3 hypotheses: Forms carbonium ions that attack DNA; acts as purine analogue and inhibits DNA and RNA synth.; interacts with SH groups | Anaphylaxis is; fever; hyper eosinophilia; allergic hepatitis; bms; GI | Rash; urticaria; photosensitivity; acn; hyper pigm | | [29, 30] |
| Temozolomide (Temodar®, Temodal®) | Brain; melanoma | Alkylates DNA at N-7 or O-6 of guanine | Anaphylaxis; fever; peripheral oedema; anaemia; neutropenia | | | [31, 32] |
| ThioTEPA (Thioplex®) | Ovarian; breast; CNS; bladder; leukemia; MM | Ethylimine cross-links DNA | Fever; anaphylaxis; wheeze; bms; thrombocytopenia; anaemia; leucopenia | | | [21, 34] |
| Antimetabolites | | | | | | |
| Pyrimidine analogues | | | | | | |
| Capecitabine (Xeloda®) | Colorectal; pancreatic; metastatic breast; stomach | Prodrug converted to 5-fluorouracil (see below) | Lymphopenia; thrombocytopenia; neutropenia; anaemia; CV symptoms; GI | | | [21, 35, 36] |

Table 2 Classification of non-targeted chemotherapeutic drugs used for cancers and their hypersensitivity/adverse reactions
| Drug generic and trade names and classification1 on the basis of mechanism | Cancer indications | Mechanism(s)1 of action | Hypersensitivity2/adverse3 reactions | References |
|---|---|---|---|---|
| Cytarabine (Cytosar-U®, Tamohine PIS®, AraC®) | AML; NHL; ALL | Cytosine arabinoside similar to deoxycytosine—competitively incorporated into DNA | Anaphylaxis; thrombocytopenia; leucopenia; fever; dyspnea; ards; GI; hepatotoxicity; bleding | [19, 21, 24, 37] |
| 5-Fluorouracil (Efudex®, Adrucil®) | Colorectal; breast; pancreatic | Inhibition of thymidyate synthase | Anaphylaxis; myelosuppression; GI; mucositis | [19, 21, 38] |
| Gemcitabine (Gemzar®) | NSCL; pancreatic; breast; ovarian; bladder | Nucleoside analogue—replaces cytidine during DNA replication | Fever; hypersens pneumonitis; fl-like symptoms; GI; leukopenia; neutropenia; anemia; thrombocytopenia | [19, 39, 40] |
| Purines | Cladribine (Link®, Movecro®) | HSL; pediatric AML | Inhibits adenosine deaminase and processing of DNA | Fever; DIHA13; increased risk virus (e.g. herpes) infection; fall in blood count; periph neuropathy; GT; dis; id | [21, 41, 42] |
| Fludarabine (Fludara®) | CLL; AML; NHL | Inhibits ribonucleotide reductase and DNA polymerase | Lymphopenia; DIHA13; thrombocytopenia; neutropenia; ahs; GI; ards | [21, 43] |
| 6-Mercaptopurine (Purinethol®) | ALL; pediatric NHL | Inhibits purine nucleotide synthesis and metabolism | Myelosuppression—toxic to bone marrow; pancreatitis; fever; GI | [44] |
| Pentostatin14,15 (Nipent®) | HSL; CLL | Inhibits adenosine deaminase and processing of DNA | Anaphylaxis; fever; flushing16; GI; leuopenia; thrombocytopenia; anemia | [21, 45, 46] |
| Folate antagonists | Methotrexate17 (formerly Amethopterin) (Trexall®) | Breast; head; neck; lung; bladder; leukemia; lymphoma | Inhibits dihydrofolate reductase | Anaphylaxis; bronchospasm; pulm infiltrates; haemolysis; anemia; agranulocytosis; myelosuppression; hepatotoxicity; GT17 | [19, 21, 24, 38, 47] |
| Pemetrexed (Alimta®) | NSCL; mesothelioma | Inhibits enzymes in purine and pyrimidine synthesis | Neutropenia; thrombocytopenia; anemia; interstitial pneum; dyspnea; GI; mucositis | [48, 49] |
| Mitotic inhibitors | Doxorubicin (Taxotere®) | Breast; NSCL; ovarian; prostate | Binds to microtubules and inhibits mitotic cell division | Anemia; neutropenia; leukopenia; bronchospasm; dyspnea; back pain | [19, 21, 24] |
| Paclitaxel (Abraxane®, Taxol®) | Lung; breast; ovarian; head and neck; bladder; testicular; Kaposi’s | Binds to β-tubulin subunits; suppresses microtubule function; blocks mitosis | bms; neutropenia; hypersens pneumonitis; dyspnea; back pain; GI; neurotox19 | [19, 21, 24] |
| Vinca alkaloids | (a) Vinblastine, (b) vincristine, (c) vindesine, (d) vinorelbine (Velban®, Oncovin®, Eldisine®, Navelbine®) | (a) HL; NSCL; head and neck; NHL; breast. (b) HL; WT; leukemia. (c) Melanoma; lung; breast; leukemia. (d) Breast; bone; NSCL | Binds to tubulin, prevents spindle formation and cell division | Anaphylaxis; fever; bronchospasm; arf; pulm oedema; pleural effusion; interstitial pneumonitis | [19, 21, 24] |

References: [19, 21, 24, 37]
### Table 2 (continued)

| Drug generic and trade names and classification¹ on the basis of mechanism | Cancer indications | Mechanism(s)² of action | Hypersensitivity³/adverse³ reactions | References |
|---|---|---|---|---|
| **Drugs that interact with or otherwise directly affect DNA²⁰** | | | | |
| **Inhibitors of DNA synthesis** | | | | |
| Hydroxyurea (Hydrea®, Droxia®) | CML; myeloproliferative disorders | Prevents reduction of ribonucleotides to deoxyribonucleotides²¹ | Fever; hypersens pneumonitis; bms; GI | [21, 53, 54] |
| DNA cross-linkers | | | | |
| Carboplatin (Paraplatin®, Paraplatin-AQ®) | Ovarian; lung; breast; head and neck; testicular; brain (children) | Cross-links DNA following formation of an aqua ligand (Cl displaced by H₂O) which binds to bases, preferably guanine | Anaphylaxis; bms; GI; bronchospasm; dyspnea; periph neuropathy | [19, 21, 55–57] |
| Cisplatin (Platinol®) | Sarcomas; ovarian; lymphomas; lung; germ cell | As for carboplatin | Anaphylaxis; bronchospasm; dyspnea; haemolytic anaemia; sens neuropathy; GI; renal tox | [19, 21, 24, 57] |
| Oxaliplatin (Eloxatin®) | Colorectal²²; ovarian; gastric | As for carboplatin | Anaphylaxis; fever; dyspnea; wheezing; sens neuropathy; GI; Types II and III hypersens²³ | [21, 24, 57, 58] |
| Mitomycin C²⁵ (Mutamycin®, Mitozyn®) | Oesophageal; bladder; breast; stomach; pancreas | Cross-links DNA by two N-alkylations of guanosine nucleoside. Specific for 5′-CpG-3′. Inhibit of thioredoxin reductase | Bone marrow toxicity; id; cosinophilic cystitis; has | [19, 21, 59] |
| Cross-linkers of opposite DNA strands | | | | |
| Pyrolobenzodiazepine dimers (e.g. SJG-136, NSC 694501)²⁹ | Ovarian²⁷ | In minor groove of DNA, the two imines react with and link N-2 positions on guanines on opposite DNA strands | Thrombocytopenia; vs (oedema, dyspnea, hypoualbinemia, pleural effusion, ascites); dlh | [60, 61] |
| Topoisomerase inhibitors | | | | |
| Topoisomerase I inhibitors | | | | |
| Topotecan (Hycautin®) | Ovarian; cervical; SCL | Stabilizes topo-I complexes inhibiting re-ligation of single strand breaks and leading to lethal double-strand breaks | bms; alveolar damage; cop; GI | [62, 63] |
| Irinotecan (Camptosar®) | Colon | Produces. Metabolite (SN-38) inhibits topo-I leading to DNA breaks | Anaphylaxis; neutropenia; anaemia; dyspnea | [64, 65] |
| Topoisomerase II inhibitors | | | | |
| Daunorubicin (Cerubidine®) | AML; ALL; neuroblastoma | Intercalates between base pairs in DNA helix (see below). Inhibits topo-II and prevents ligation of nucleotide strand after double strand breaks | bms; anaphylaxis; fever; cardiac tox; GI | [19, 21, 24, 66] |
| Doxorubicin (Doxil®) | HL; haematological; ovarian; breast; lung; bladder; Kaposis | As for daunorubicin | Anaphylaxis; myelosuppression; bronchospasm; hir; cardiotoxicity; GI | [19, 21, 24, 67] |
Table 2 (continued)

| Drug generic and trade names and classification¹ on the basis of mechanism | Cancer indications | Mechanism(s)² of action | Hypersensitivity³, adverse³ reactions | Systemic | Cutaneous | References |
|---|---|---|---|---|---|---|
| Etoposide (Etopophos®) | SCL; germ cell; Kaposi’s and ES | Inhibits topo-II and ligation of cleaved DNA leading to single and double-stranded DNA breaks | Myelosuppression; fever; hypotension; tachycardia; bronchospasm; dyspnea; GI bms; fever; hypotension; bronchospasm; wheeze; intravascular haemolysis³⁰; GI | Flashing; exanthema; urticaria; alopecia; puritus; hyper pigm | | [19, 21, 24] |
| Teniposide (Vumon®, Vehem®) | ALL | Complexes with topo-II and DNA causing single and double-stranded breaks in DNA and inhibition of strand re-ligation | Stomatitis; alopecia; rash; facial oedema; urticaria³⁰ | | | [19, 21, 24] |
| Mitoxantrone (Novantrone®) | Breast; AML; NHL; ALL | Intercalates with DNA and inhibits topo-II disrupting DNA repair and synthesis | Anaphylaxis; myelosuppression; GI; CV effects | isr; rash; purpura; nail discoloration; stomatitis; alopecia; drees | | [21, 68] |
| Intercalation with DNA | | | | | | |
| Actinomycin D³¹ (Cosmegen®) | WT; ES; testicular; trophoblastic | Intercalates between adjacent base pairs of DNA; prevents elongation of RNA by RNA polymerase; stabilizes complexes of topo-I and topo-II with DNA | Leukopenia; anaemia; GI; hepatotoxic effects | Rash; isr; stomatitis; rrr; alopecia; photosens | | [69–71] |
| Daunorubicin | AML; ALL; neuroblastoma | Intercalates between base pairs in DNA helix preventing DNA replication and protein synthesis | bms; anaphylaxis; fever; cardiac toxic; GI | Rash; uricemia; angioedema; hyper pigm | | [19, 21, 24, 66] |
| Doxorubicin | HL; haematological; ovarian; breast; lung; bladder; Kaposi’s | See also topoisomerase II inhibitors (above) | Anaphylaxis; myelosuppression; bronchospasm; hir; cardiotoxicity; GI | Rash; acnl erythema; isr; puritus; urticaria; angioedema; neph; rrr | | [19, 21, 24, 67] |
| Inducers of DNA breaks | | | | | | |
| Bleomycin³² (Blenoxane®) | HL; NHL; SCC; testicular; squamous cell | Causes breaks in DNA by unresolved mechanism (may chelate metal ions producing superoxide and hydroxide radicals; blocks uptake of thymidine by DNA). Mediates oxidative degradation of cellular RNA | Hypotension; intermit pneumonitis; pulm fibrosis; fever³³ | Rash; pruritus; angioedema; scleroderma-like; Raynaud’s syndrome; sj; alopecia; erythema; hyper pigm | | [19, 21, 72] |
| Inhibitors of DNA methylation | | | | | | |
| Azacitidine (Vidaza®) | MDS; AML | Inhibition of DNA methyltransferase which prevents DNA synthesis; cytotoxic to abnormal haematopoietic cells in bone marrow via incorporation into DNA and RNA | bms; thrombocytopenia; anaemia; neutropenia; GI pyrexia; periph oedema; dyspnea; cough; anaphylaxis | Rash; isr; ecchymoses; xerosis; peetechia; erythema; urticaria; pruritus; Sweet’s syndrome | | [73, 74] |
| Inhibitors of DNA repair | | | | | | |
| PARP inhibitors (e.g. Iniparib, Olaparib, Niraparib, Rucaparib, Veliparib) | In clinical trials for breast, lung and ovarian cancers and metastatic melanoma | Block repair of DNA single strand breaks (ssb) trap PARPs 1 and 2 at damaged DNA sites—trapped DNA-PARP complexes more cytotoxic than un repaired ssb | Myelosuppression | Drugs still in clinical trials with little definite information on adverse reactions available | | [75, 76] |
| Disruption of protein synthesis | | | | | | |
| L-Asparaginase³⁴ (Elspar®) | ALL; AML | Hydrolyses l-asparagine to l-aspartic acid and NH₃. Some leukemia cells cannot synthesize l-asparagine resulting in inhibition of protein synthesis | Anaphylaxis³⁵; laryngospasms; bms (transient); GI; pancreatitis | Urticaria; angioedema; msh; pruritus; stomatitis; ten | | [19, 21, 24] |
| Drug generic and trade names and classification on the basis of mechanism | Cancer indications | Mechanism(s)\(^1\) of action | Hypersensitivity\(^2\)/adverse\(^3\) reactions | References |
|---|---|---|---|---|
| **Systemic** | | | | |
| Lenalidomide (Revlade\(^®\)) | MM; MDS | Multiple actions: Inhibits proliferation and apoptosis of tumour cells; activates immune T and NK cells; induces cytokine production; anti-angiogenic property | Thrombocytopenia; anaemia; neutropenia; hypersensitivity pneumonitis; thrombosis; rash (morbilliform, urticarial, dermatitic), pruritus; SJS; erythema; ecchymosis; erythema multiforme | [21, 77] |

\(^1\) Some drugs exert their action by more than one mechanism and might therefore be classified into more than one category. The classification shown is deemed to be the most appropriate one.

\(^2\) Reactions such as weakness, fatigue, headache etc. and symptoms like nausea, vomiting, diarrhea, constipation, appetite suppression etc. are not listed since they are common to so many of the drugs. Where a latter symptom(s) (i.e. nausea etc.) is important, it is referred to as ‘GI’

\(^3\) Allergic reactions ~1%

\(^4\) Antibiotic from *Streptomyces achromogenes*

\(^5\) Streptomicin toxic to pancreas β cells

\(^6\) Twenty percent patients with metastatic malignant melanoma are hypersensitive

\(^7\) Hepatic veno-occlusive disease (Budd–Chiari syndrome)

\(^8\) Precise mechanism not known

\(^9\) Approximately 28% of population dihydropyrimidine dehydrogenase deficient, i.e. genetic inability to metabolize pyrimidine-based drugs

\(^10\) Up to 73% patients have rash. Type I reaction rare

\(^11\) Palmar–plantar dermatitis a risk factor for hypersensitivity reactions

\(^12\) Drug may induce autoantibodies without becoming bound to red cells

\(^13\) Antibiotic from *Streptomyces venaticus*

\(^14\) Concomitant use of allopurinol a risk of hypersensitivity reactions

\(^15\) Incidence of hypersensitivity 0.5–1%

\(^16\) Implicated in type I (urticaria, angioedema), type II (haemolytic anaemia) and type III (pneumonitis, cutaneous vasculitis) hypersensitivity reactions

\(^17\) Incidence of cutaneous reactions ~14%

\(^18\) Reaction 10–16% without premedication

\(^19\) Inhibits DNA synthesis/repair or methylates or damages DNA

\(^20\) Acts in S phase but detailed mechanism of action remains uncertain

\(^21\) Cutaneous manifestations in up to 35% patients

\(^22\) Administered in combination with 5-fluorouracil, folinic acid and leucovorin—FOLFOX

\(^23\) Type II thrombocytopenia and type III immune complex-mediated urticaria hypersensitivities occur

\(^24\) Antibiotic from *Streptomyces coelicoflavus* or *Streptomyces lividans*

\(^25\) Cutaneous reactions in 9% patients after intravesical administration

\(^26\) Now in phase 2 trial

\(^27\) Insufficient data available yet

\(^28\) In development

\(^29\) Reactions in 41% children with ALL treated with teniposide and other chemotherapy drugs

\(^30\) Antibiotic from *Streptomyces parvulus*. Also known as dactinomycin

\(^31\) Glycopeptide antibiotic from *Streptomyces verticillus*

\(^32\) Fever in up to 25% patients and fulminant reactions in up to 8%

\(^33\) Enzyme from *Escherichia coli* or *Erwinia carotovora*

\(^34\) Frequency of antibodies to L-asparaginase up to 79% in adults and 70% in children. Hypersensitivity to *E. coli* L-asparaginase up to 45%. *Erwinia* enzyme used in patients allergic to *E. coli* enzyme
tamoxifen, aromatase inhibitors such as exemestane [136] and anastrozole that prevent the conversion of androgens to oestrogens and some hormone analogues (goserelin) [137] and anti-androgens (bicalutamide, flutamide) have been part of the targeted therapeutic approach to breast and prostate cancers [138] (Table 3). The first signalling proteins to be utilized as targets for a new generation of unique anti-cancer drugs were protein kinases. The strategy adopted was based on the instability of the cancer genome compared to the normal cell. The Philadelphia translocation t(9;22)(q34;q11) is a chromosomal defect resulting in gene fusion of the BCR gene on chromosome 22 (region q11) and ABL tyrosine kinase gene on chromosome 9 (region q34), producing the BCR-ABL oncogene. The Philadelphia chromosome abnormality is seen in 95 % of chronic myeloid leukemia (CML) patients and 15 to 30 % of adults with acute lymphoblastic leukemia (ALL). CML cells show absolute dependence (‘oncogene addiction’) on the kinase activity of protein Bcr-Abl [139], and this dependence was first exploited by the development of the drug imatinib (Gleevec®) which inhibits the tyrosine kinase. In CML cases resistant to imatinib, broader spectrum tyrosine kinases such as dasatinib (Sprycel®), which blocks both Bcr-Abl and Src tyrosine kinases, may be used [140] (Table 3). Nilotinib is another protein tyrosine kinase inhibitor targeting Bcr-Abl. Inhibitors of receptor tyrosine kinases targeting epidermal growth factor receptor (EGFR; ErbB1; HER1; a member of the ErbB family of receptors), vascular endothelial growth factor receptors (VEGFRs) and platelet-derived growth factor receptors (PDGFRs) have also found use in the clinic as effective targeted anti-tumour drugs [96]. Some receptor tyrosine kinase inhibitors include gefitinib and erlotinib (both inhibitors of EGFR), lapatinib (inhibits ErbB1 and ErbB2), vatalanib (VEGFR-1 and VEGFR-2), sorafenib (VEGFR, PDGFR and c-Kit) and sunitinib (VEGFR, PDGFR and c-Kit). Other targeting strategies summarized in Table 3 that interfere with signal transduction and which have entered, or are about to enter the clinic, include the more recently introduced mTOR serine/threonine kinase inhibitors that target the Raptor complex [141] (drug examples include everolimus and temsirolimus), histone deacetylase (HDAC) inhibitors (e.g. romidepsin and vorinostat) that arrest the cell cycle [142], targeting the PML-RARα oncoprotein (arsenic trioxide) [143] and drugs (e.g. bexarotene) binding to retinoid receptors [144]. Still other targeting strategies are represented by pralatrexate, a folate analogue that accumulates in cancer cells over-expressing protein RFC-1 [145] and the cytotoxic chimeric proteins denileukin difitox [146] and ziv-afibercept [147, 148] (Table 3). Finally, proteasome inhibitors such as bortezomib and carfilzomib work against the cells of multiple myeloma and mantle cell lymphomas in perhaps the most fascinating of all the current targeted mechanisms [149, 150]. These drugs inhibit proteasomes by binding to proteolytic catalytic sites in the 20S proteasome core (Fig. 2). This is thought to prevent degradation of pro-apoptotic factors permitting killing of cancer cells.

4 Side effects of chemotherapeutic drugs

Adverse reactions to important examples from the main groups of non-targeted and molecularly targeted agents are numerous and often diverse, and with the large number of drugs currently employed in cancer therapy, the task of conveying a succinct, informative and accurate picture of adverse reactions at the level of both individual drugs and different categories of drugs with common features is a formidable one. From information obtained from the published medical/scientific literature, research studies and FDA and drug company data, Table 2 summarizes the most important systemic and cutaneous adverse reactions of the non-targeted anti-cancer drugs. With the exception of only one drug, lenalidomide which has multiple actions, the other drugs have been arranged in different categories based on common mechanisms of action. It has been claimed that the problem of hypersensitivity reactions to these chemotherapeutic drugs is often not reported by oncologists and therefore underestimated and that they can be divided into three groups with high, intermediate or low potential for causing hypersensitivity reactions [24]. Drugs included in the high group are platinum compounds, taxanes, epipodophyllotoxin topoisomerase II inhibitors, L-asparaginase and procarbazine. The anthracyclines (daunorubicin, doxorubicin and epirubicin), 6-mercaptopurine and methotrexate are in the intermediate division while cytarabine, cyclophosphamide and ifosfamide make up the low potential group. Antimetabolites such as capecitabine, gemcitabine and 5-fluorouracil commonly provoke acral erythema syndrome (Fig. 3) [153], and in common with cyclophosphamide and doxorubicin, hyperpigmentation may occur. Mitotic inhibitors, the taxanes and Vinca alkaloids, often produce alopecia and nail abnormalities and radiation recall reactions are seen (Table 2). With some drugs such as the PARP inhibitors that are still in clinical trials, little information is available yet on adverse, let alone hypersensitivity, reactions (Table 2).

The inclusion of reactions to 16 kinase inhibiting drugs demonstrates the current importance of tyrosine and serine/threonine kinase inhibitors for the targeted treatment of an expanding range of tumours (Table 3). While the aim of signal transduction therapy is to kill the cancer cells with minimal collateral damage (Fig. 1), even a quick glance at the catalogue of side effects in the table shows that, just as with the non-targeted drugs (but less so), cutaneous, gastrointestinal and haematopoietic cells are often still affected. Cytotoxic effects such as anaemia, thrombocytopenia and neutropenia occur less often and usually with less severity than with, say, antimetabolites and alkylating agents, but a number of the...
Table 3  Classification of targeted chemotherapeutic drugs used for cancers and their hypersensitivity/adverse reactions

| Drug generic and trade names and classification on the basis of mechanism | Cancer indications | Mechanism(s) of action | Hypersensitivity/adverse reactions | References |
|---|---|---|---|---|
| **Signal transduction inhibitors** | | | | |
| **Non-receptor tyrosine kinase inhibitors** | | | | |
| Bosutinib (Bosulif®) | Ph+CML | Targets Abl and Src kinases inhibiting autophosphorylation of both | Anaphylaxis; thrombo; neutro; anaemia; oedema; hepatotoxic; pneumonia; pyrexia; cough | Rash; pruritus | [78, 79] |
| Crizotinib (Xalkori®) | NSCL (Trials: ALCL; neuroblastoma; solid tumours) | Inhibits tyk kinase activity of fusion protein EML4-ALK | Pneumonitis; oedema; neuropathy; arthralgia; vision disorders; cough; dyspnea; QT prolongation; GI | Rash; pruritus; stomatitis | [80, 81] |
| Dasatinib (Sprycel®) | CML; Ph+ ALL | Inhibits Bcr-Abl, Src, c-Ki6, ephrin (EphA2) and other tyk kinases (PDGFRβ) | Abnormal platelets; neutro; anaemia; peripheral oedema; pleural effusion; palsy; GI; CV; QT prolongation; dyspnea | Rash; pruritus; nail disorders; pigm; panniculitis; dermatitis | [82, 83] |
| Imatinib (Gleevec®) | Ph+CML; Ph+ALL; GI; stromal; some CEL | Blocks Bcr-Abl tyk kinase from phosphorylating | Thrombo; neutro; anaemia; oedema; hypersens pneum; hepatotoxic; tks; CV; GI | Rash; pruritus; angioedema; vasculitis; hyper-pigm; dress; agep; sjs erythema multiforme | [84–86] |
| Nilotinib (Tasigna®) | CML | Inhibits Bcr-Abl, c-Ki6, EphA3, EphA5, PDGFRβ and other tyk kinases | Thrombo; neutro; leuko; anaemia; oedema; hepatotoxic; GI; pneumonia; dyspnea; QT prolongation | Rash; pruritus; erythema; eczema; urticaria; alopecia; xerosis; ecchymosis | [87] |
| Ponatinib (Iclusig®) | CML; Ph+ALL | Targets pan Bcr-Abl | Thrombo; neutro; anaemia; leuko; thrombosis; hepatotoxic; CV; tks; pancreatitis; GI; arthralgia/myalgia | Rash; xerosis | [88] |
| **Receptor tyrosine kinase inhibitors** | | | | |
| Erlotinib (Tarceva®) | NSCL; pancreatic | EGFR inhibitor—binds to ATP binding site of receptor interrupting signal cascade | Ild; hepatotoxic; thrombo; anaemia; GI; renal disorders | Rash; pruritus; ppr; xerosis; hyper-pigm; hfsr; digital fissures; alopecia; paronychia; eyelid/brow changes; bullous reactions | [89, 90] |
| Gefitinib (Iressa®) | NSCL; breast | EGFR inhibitor | GI; ild; renal/urinary disorders; haemorrhage; | hfsr; pp; pruritus; ocular disorders; nail and hair disorders; stomatitis | [90, 91] |
| Lapatinib (Tykerb®) | Breast; other solid tumours | Inhibits tyk kinase activity of HER2 and EGFR | Anaphylaxis; cardiac tox; hepatotoxic; ild; GI; QT prolongation | Rash; hfsr; pruritus; xerosis; nail disorders; paronychia; alopecia | [92] |
| Pazopanib (Votrient®) | RCC; STS; (ovarian; NSCL?) | Inhibits several tyk kinases incl VEGFRs, PDGFRs, FGFR2, c-Kit | Hepatotoxic; hypertension; GI; QT prolongation; haemorrhage; hypothyroidism; rpls | hfsr; hair/skin depigm; exfol rash; nail disorders; xerosis; stomatitis; alopecia; mucositis | [93] |
| Regorafenib (Stivarga®) | Colorectal; GIST | Inhibits multi kinases incl VEGFRs 2 and 3, PDGFR, c-Ki6, RET, RAF, TIE2 | Hepatotoxic; hypertension; haemorrhage; GI; dysphonia | hfsr; rash; xerosis; mucositis; desquamation; pruritus | [94, 95] |
| Sorafenib ( Nexavar®) | RCC; HCC | Inhibits VEGFR2 and PDGFRβ signalling blocking tumour angiogenesis. Blocks RAF kinase involved in cell growth and proliferation | Haemorrhage; cardiac tox; hypertension; GI; flushing; thrombo; neutro; anaemia; leuco; lympho; rpls | hfsr; rash; desquamation; erythema; xerosis; pruritus; alopecia; sjs; facial acne; stomatitis; bullous eruptions | [96–99] |
| Sunitinib (Sutent®) | RCC; GIST; PNET | Inhibits VEGFR2, PDGFR3 and c-Ki6, inhibiting angiogenesis and cell proliferation. Inhibits phosphorylation of FLT3 | Hepatotoxic; hypertension; CV; GI; QT prolongation; haemorrhage; hypothyroidism; fever; periph oedema; dyspnea; cough | hfsr11; yellow skin; rash; xerosis; hair depigm; alopecia; erythema; pruritus; stomatitis | [96, 98–100] |
| Vandetanib (Caprelsa®) | MTC | Inhibits cell receptor tyk kinases VEGFR, EGFR and RET | Hypertension; cardiac tox; QT prolongation; haemorrhage; ild; rpls; cough | Rash and acne (? ppr); xerosis; pruritus; dermatitis; hfsr; erythema; photosens; sjs | [101] |
| Drug generic and trade names and classification¹ on the basis of mechanism | Cancer indications | Mechanism(s)¹ of action | Hypersensitivity²/adverse³ reactions | References |
|---|---|---|---|---|
| **mTOR serine/threonine kinase inhibitors** | | | | |
| Everolimus (Afinitor®) | RCC; breast; PNET; SEGA | Binds FK-binding protein FKB P12 and drug-protein complex inhibits mTOR. This blocks binding of regulatory-assoc protein of mTOR causing dephosphorylation of S6K1 and apoptosis of cancer cells | Pneumonitis infections; ↓ platelets, neutrophils, Hb and lymphocytes; renal failure; oedema; oral ulcers; GI; fever; dyspnea | Acneform rash (? ppr); erythema; stomatitis; nail disorders; xerosis [102] |
| Temsirolimus (Torisel®) | RCC | Binds FK-binding protein FKB P12 and drug-protein complex inhibits mTOR. This blocks binding of regulatory-assoc protein of mTOR causing dephosphorylation of S6K1 and apoptosis of cancer cells. Also inhibits tumour angiogenesis by reduced synthesis of VEGF | hi¹², ikl; renal failure; thrombo; leuko; anaemia; oedema; GI; bowel perforation; eye disorders¹³ | Rash (mpr, exfoliative dermatitis); nail disorders; pruritus; xerosis; acne (? ppr); stomatitis [103] |
| **Histone deacetylase inhibitors** | | | | |
| Romidepsin (Isotadex®) | CTCL | Inhibits histone deacetylase altering gene expression and causing cell cycle arrest and apoptosis | Thrombo; neutro; lympho; anaemia; QT prolongation; GI; dysgeusia; pyrexia | Exfol dermatitis; itching; pale skin [104] |
| Vorinostat (Zolinza®) | CTCL | Induces growth arrest, differentiation or apoptosis due to accumulation of acetylated proteins incl BCL6, p53 and Hsp90¹⁴ | Thrombo; anaemia; pulm embolism; thrombosis; GI; dysgeusia | Exfol dermatitis; pruritus; alopecia [105, 106] |
| **Drugs that target the PML-RARα oncoprotein** | | | | |
| Tretinoin (ATRA) (Vesanoid®) | APL | APL results from expression by NB4 cells of PML-RARα oncoprotein which increases cell surface levels of plasminogen receptor S100A10 and fibrinolytic activity. ATRA down regulates S100A10 blocking fibrinolytic activity (and hence haemorrhage) | Retinoic acid syndrome (dyspnea, fever, periph oedema, weight gain); els; retinoid tox; respiratory and ear disorders | Erythema; xeroderma; photosens; ich; increased risk sunburn [107] |
| Arsenic trioxide (Trisenox®) | APL | As₂O₃ binds to oncogenic fusion protein PML-RARα, induces increased interaction with a ubiquitin-like protein-modifier SUMO-conjugating enzyme UBC9, enhanced SUMOylation and degradation | Thrombo; neutro; hyperleukocytosis; anaemia; hypertension; vasculitis; differentiation syndrome; dyspnea; QT prolongation; GI | Rash; erythema; pruritus; facial oedema; ecchymosis [108] |
| **Drugs binding to retinoid receptors** | | | | |
| Bexarotene (Targretin®) | CTCL | Synthetic retinoic acid deriv. Binds and activates retinoid X receptors inducing changes in gene expression, decreased cell proliferation, apoptosis of some cancer cell types and tumour regression | Lipid abnormalities; pancreatitis; leuko; anaemia; periph oedema; GI; hypothyroidism; infection; chills; fever | ppr; mpr; acne; vesicular bullous rash; keratitis; conjunctivitis; corneal lesions; alopecia; skin nodules; photosens; xerosis [109] |
| **Proteasome inhibitors** | MM; MCL; NHL | Inhibits proteasomes by binding via its β atom to β-subunit chymotrypsin and caspase catalytic sites—may suppress degradation of pro-apoptotic factors allowing for killing of neoplastic cells | Periph neuropathy; GI; thrombo; neutro; leuko; anaemia; hypotension; cardiac disorders; ards; rpls; ifs; hepatic events; pyrexia; dyspnea; pneumonia | Rash¹¹; pruritus; erythema; urticaria; facial oedema; eczema; vasculitis; ten; Sweet’s-like syndrome [110–112] |
| Drug generic and trade names and classification | Cancer indications | Mechanism(s) of action | Hypersensitivity adverse reactions | References |
|-----------------------------------------------|--------------------|------------------------|-----------------------------------|------------|
| Carfilzomib (Kyprolis®) MM                    | Cancer             | Binds to and inhibits chymotrypsin-like activity of the 20S proteasome | Cardiac disorders; pulm hypertension; dyspnea; infusion reactions; tls; thrombo; anaemia; neutro; hepatotox; periph neuropathy | [113] |

### Hormones, hormone analogues and hormone antagonists

#### Inhibitors of hormone synthesis

Hormone receptor antagonists

**Selective oestrogen receptor modulators (SERMS)**

- **Tamoxifen**
  
  Breast (oestrogen receptor-positive) and post-menopausal women
  
  Competitive antagonist of oestrogen receptor, inhibiting breast cancer cells requiring oestrogen to grow; anti-angiogenic effect
  
  Hot flashes; vaginal discharge; endometriosis; reduced vision; thrombosis; interstis pneumonitis
  
  Pruritus vulvae; urticaria; angioedema; vasculitis; erythema multiforme; sjs
  
  [124, 125]

### Targeted anti-folate drug

- **Pralatrexate (Folotyn®)**
  
  PTCL
  
  Folate analogue antimetabolite accumulates in cancer cells which overexpress the protein RFC-1 and interferes with DNA synthesis and leading to cell death
  
  Thrombo; febrile neutro; anaemia; mucositis; pyrexia; dyspnea; foetal harm; tls
  
  Rash; pruritus; exfoliation; ulceration; ten
  
  [127]
| Drug generic and trade names and classification | Cancer indications | Mechanism(s) of action | Hypersensitivity/adverse reactions | References |
|-----------------------------------------------|--------------------|------------------------|-----------------------------------|------------|
| **Targeted cytotoxic chimeric proteins**      |                    |                        |                                   |            |
| Denileukin diftitox (Ontak®)                  | CTCL               | Modified diphtheria toxin and IL-2 receptor fused to give recombinant toxin. Toxin released intracellularly causes ADP-ribosetransferase-mediated inhibition of protein synthesis and cell death | Infusion reactions; pyrexia; cns; vision disturbances; periph oedema; dyspnea; cough; GI | [128–130] |
| Ziv-Aflibercept (Zaltrap®)                    | MCRC               | Fusion protein of Fc portion of IgG and ligand-binding domains of VEGFR1 and VEGFR2. Acts as a soluble decoy VEGF receptor and angiogenesis inhibitor | leuko; neutro; thrombo; haemorrhage; GI perforation; proteinuria; hypertension; venous thromboembolism | [131] |

**Table 3 (continued)**

* Some drugs exert their action(s) by more than one mechanism and might therefore be classified into more than one category. The classification shown is deemed to be the most appropriate one.

* Reactions known, or suspected, of having an immunological basis

* Reactions such as weakness, fatigue, headache etc. and GI symptoms like nausea, vomiting, diarrhoea, constipation, appetite reduction, dyspepsia etc. are not recorded here since they are common to so many of the drugs. Where such reactions are important, they are referred to collectively as 'GI'

* Proto-oncogene c-Kit, mast/stem cell growth factor receptor. Also called tyrosine-protein kinase kit or CD117

* Belongs to ephrin receptor family of tyrosine kinases

* Ponatinib used to treat refractory CML including patients with T3151 ‘gatekeeper’ mutation

* The enzyme RAF proto-oncogene ser/thre-protein kinase (also called c-Raf or Raf-1). Functions in MAPK/ERK signal transduction pathway

* TIE2, angiopoietin receptor that binds angiopoietin growth factor required for angiogenesis

* Thirty to 60 % of patients

* FLT3, CD117; Fms-like tyrosine kinase; receptor-type tyrosine protein kinase FLT3

* Fifteen to 20 % of patients

* May occur on first and subsequent infusions. Reactions include flushing, chest pain, dyspnea, hypotension, anphoea and anaphylaxis. Allergic/hypersensitivity reactions occur in ~9 % of patients (FDA)

* Seven percent of patients

* BCL6 protein—zinc finger transcription factor, a sequence-specific repressor of transcription. p53 is a tumour suppressor protein that regulates the cell cycle. Hsp90 is a molecular chaperone assisting protein folding and stabilizes some proteins in tumour growth

* Incidence 8–18 %

* Skin and subcutaneous tissue adverse events in phase 2 trial on 526 patients with MM=36 %. Discontinuations due to these adverse events=1 %

* Also called lateinizing hormone

* In 2006, NCI stated that raloxifine was as effective as tamoxifen in reducing the incidence of breast cancer in post-menopausal women at increased risk

* Nineteen percent of treated patients experience skin reactions
targeted agents show their own fairly unusual effects including, for example, a lengthened QT interval, a hand–foot skin reaction (Fig. 4) ([151]; see later) significantly different from the well-known drug-induced acral erythema (compare Figs. 3 and 4) and a characteristic papulopustular rash (Fig. 5) [154]. Inhibition of the EGFR in skin often produces xerosis, skin fissures, nail alterations, paronychia and periungual ulcers. Multikinase inhibitors like sorafenib [151] and sunitinib frequently exhibit skin toxicity and produce fever, oedema and occasionally inflammatory actinic keratosis and bullous manifestations (Table 3).

5 Hypersensitivity reactions of chemotherapeutic drugs

Amongst anti-neoplastic drugs, methotrexate and oxaliplatin, for example, can provoke types I, II and III hypersensitivities. Some drugs and what at first sight appear to be ‘allergic’ drug reactions do not fit easily into the four Gell and Coombs categories. As mentioned, some reactions to contrast media and NSAIDs are two examples [8, 9] that readily spring to mind as well as skin reactions such as alopecia, folliculitis and hyperpigmentation. The situation where the mechanism of a drug-induced adverse response remains un-investigated or poorly understood is often seen in reactions to chemotherapy drugs where the haematological system, liver, lungs and skin are frequently affected (Tables 2 and 3). It has been said that so-called idiosyncratic drug reactions most often manifest as cutaneous reactions, haemato- and hepato-toxicities and less frequently as anaphylaxis and autoimmunity, but some investigators believe that most idiosyncratic reactions are hypersensitivities occurring via an immune-mediated response [156]. Cytotoxic chemotherapy typically suppresses haematopoiesis causing thrombocytopenia, neutropenia and anaemia, and drug-induced hepatotoxicity is a well-known cause of liver injury. However, all four conditions can be instigated by drugs via an immune mechanism, and this possible difference in the underlying mechanisms of these most
serious drug side effects frequently remains unexplored. In fact, autoimmune forms of haemolytic anaemia and thrombocytopenia are well-known complications of lymphoproliferative diseases and occur predominantly in patients with lymphocytic leukemias [157]. Specific aspects, including the special relevance and significance of chemotherapy drug-induced thrombocytopenia, neutopenia (and agranulocytosis), anaemia, liver injury, lung disease and cutaneous reactions, are considered below.

5.1 Type I hypersensitivity reactions to chemotherapeutic drugs

Type I, IgE antibody-mediated hypersensitivity, also known as immediate, or sometimes anaphylactic hypersensitivity, can affect a single organ such as the nasopharynx (allergic rhinitis), eyes (conjunctivitis), mucosa of mouth/throat/tongue (angioedema), bronchopulmonary tissue (asthma), gastrointestinal tract (gastroenteritis) and skin (urticaria, eczema), or multiple organs (anaphylaxis), causing symptoms ranging from minor itching and inflammation to death. Symptoms of a drug-induced pseudo-allergic or anaphylactoid reaction resulting from the direct release of histamine and producing flushing, rash, pruritus, urticaria, hypotension and mucous secretion can sometimes make it difficult to distinguish these symptoms from a true, IgE-mediated type I hypersensitivity reaction. To do this, investigations designed to look for an immune basis of the reactions are necessary. Symptoms of drug-induced anaphylaxis generally include cardiovascular collapse, bronchospasm, urticaria, angioedema, erythema,
gastrointestinal symptoms (abdominal pain, vomiting, diarrhoea, haematemesis, melena) and pulmonary oedema [158, 159]. Symptoms are generally more severe in anaphylactic than in anaphylactoid reactions with cardiovascular collapse and bronchospasm occurring more frequently in the former and cutaneous manifestations seen more often in the latter [160–162]. In fact, the appearance of bronchospasm indicates that a drug reaction is more likely to be anaphylactic than anaphylactoid [163].

It is uncommon to find a drug, even those used with only moderate frequency, that has not provoked anaphylaxis, urticaria and/or angioedema in at least one rare individual so it is not surprising to see type I hypersensitivity responses well represented in the lists of adverse reactions to the many chemotherapeutic drugs. Such reactions remain rare but are seen more often with the platinum agents, l-asparaginase and methotrexate. Although there has been no clear and unequivocal demonstration of the existence of IgE antibodies to the platinum drugs, sensitizations only after multiple infusions, positive skin tests and anaphylactoid-like reactions are all taken as evidence in support of a type I hypersensitivity mechanism with these agents [164]. This conclusion is supported by positive oxaliplatin [164, 165] and carboplatin [166] skin tests that show a sensitivity of 75 to 100 %. L-Asparaginase seems to be the drug with the highest risk of provoking a type I response with an allergic reaction occurring in up to 43 % and serious anaphylaxis in a little less than 10 % of treated patients. One study employing skin and Prausnitz–Küstner tests indicated an IgE-mediated mechanism in five anaphylactic patients [167], whereas anaphylaxis to l-asparaginase in seven of eight cases was concluded to be due to complement activation induced by immune complexes of the drug with IgG and IgM antibodies [168]. Reactions are also well-known to procarbazine; the alkylating agents cyclophosphamide, melphalan, mechlorethamine, dacarbazine, temozolomide and thioTEPA; antimetabolites cytarabine and 5-fluorouracil; the Vinca alkaloid mitotic inhibitors; topoisomerase inhibitors irinotecan, daurubicin, doxorubicin and mitoxantrone; aromatase inhibitors exemestane and letrozole; some hormones and hormone analogues goserelin, leuprolide and tamoxifen; and targeted drugs such as the tyrosine kinase inhibitor bosutinib and mTOR inhibitor everolimus [18, 19, 21, 24]. Although some reactions that resemble type I reactions are dismissed as not true immediate, IgE-mediated responses because they occurred on first exposure, it should be remembered that many such drug reactions do occur on first exposure, for example, to neuromuscular blockers, contrast media, proton pump inhibitors, quinolones and others [169, 170]. Examples of some anti-cancer drugs that may provoke anaphylactoid reactions include 5-fluorouracil and the topoisomerase II inhibitors teniposide and etoposide which induce erythema, chills, fever, tachycardia, hypotension and urticaria/angioedema in up to 10 % of patients often during the first infusion.

5.2 Type II reactions

Anti-neoplastic drugs may cause a number of type II antibody-mediated cytotoxic hypersensitivity reactions, but such reactions are often not referred to by this title in the oncology and some other medical specialty literatures.
5.2.1 Drug-induced thrombocytopenia

In adults, a normal platelet count is $\sim 150–400 \times 10^3/\mu l$, and if this drops to $\sim 50 \times 10^3/\mu l$, most patients experience thrombocytopenia. For patients with drug-induced thrombocytopenia (DIT), the condition is invariably severe at counts around $20 \times 10^3/\mu l$. Many anti-neoplastic drugs such as alkylating agents, mitotic inhibitors and anti-metabolites cause a dose-dependent bone marrow suppression of platelets and all of the other haematopoietic cell lines. Some haematopoietic cells, including platelets, can be affected by drug-induced type II hypersensitivity reactions. Immune-mediated, drug-dependent destruction of platelets may occur via a number of different mechanisms, including production of a new antigen after covalent binding of drug to platelet membrane glycoprotein, binding of drug to the platelet GPIIb/IIIa complex inducing a conformational change and a new antigenic determinant, non-covalent binding to platelet membrane glycoprotein forming a drug–glycoprotein complex that reacts with antibody, binding of the Fab fragment of the chimeric human–mouse monoclonal antibody abciximab to the platelet glycoprotein receptor GPIIb/IIIa followed by subsequent recognition by antibody, production of drug-induced autoantibody that reacts with the platelet surface glycoprotein and, lastly, reaction of heparins with platelet factor 4 producing an antibody-reactive antigen, immune complexes bound to the platelet Fc receptor and platelet activation [171, 172].

Data from 2000 to 2007 on 75,243 chemotherapy regimens of associated anaemia and thrombocytopenia contained in medical records of 47,159 US patients [173] with lung, breast, ovarian, head and neck or colorectal cancers treated with platinum-, anthracycline-, gemcitabine- or taxane-based regimens were assessed in an effort to determine the prevalence of thrombocytopenia in patients undergoing chemotherapy for solid tumours. The prevalence of thrombocytopenia ranged from 21.9 % in patients treated with non-taxane-based regimens to 64.2 % in the gemcitabine-based treatment groups. During the same time, from 46.4 to 59 % of the patients developed anaemia. In a similar retrospective hospital-based cohort study from the Netherlands on the frequency and relative risk of chemotherapy-induced thrombocytopenia in 614 patients receiving one of 37 different regimens, the overall frequency of thrombocytopenia was 21.8 % [174]. The highest frequencies were seen in patients receiving carboplatin (81.8 %) and combination therapies that included carboplatin (58.2 %), gemcitabine (64.4 %) or paclitaxel (59.3 %). The overall incidence of thrombocytopenia without other cytopenias was 6.2 % although this was higher for combination therapies that included oxaliplatin (28.6 %) or gemcitabine (28.9 %). The highest relative risks of thrombocytopenia were for combination therapies of carboplatin/gemcitabine and carboplatin/paclitaxel/etoposide. These findings demonstrate that chemotherapy-induced thrombocytopenia is an important problem in clinical practice and that further research is needed to increase understanding of its clinical consequences. Importantly, some of the findings show that a number of chemotherapeutic drugs cause immune-mediated thrombocytopenia.

Based on mechanisms, there are two main categories of DIT: drug-induced marrow suppression and platelet removal by immune processes. Immune DIT generally tends to precipitate a more severe reaction than the non-immune form induced by myelosuppressive drugs, and while thrombocytopenia caused by the latter agents is often anticipated, not surprising, and even expected with some of the drugs, there will be occasions when there is a need to distinguish the two different forms. The time course of each is generally different, and this is a valuable diagnostic distinction. Non-immune marrow suppression is usually slow, often developing over weeks whereas the time for immune DIT is of the order of 1 to 2 weeks although the delay for a few patients may be only a few hours [171, 172]. Besides the mAbs rituximab and trastuzumab [175] used for cancer therapy, numerous cases involving oxaliplatin ([58, 176] and references therein), fludarabine [177], irinotecan [65], actinomycin D [178],

![Fig. 4](https://example.com/fig4.jpg)  
Hand–foot skin reaction caused by the multikinase inhibitor sorafenib (compare with Fig. 3). Hyperkeratosis, manifesting as painful yellowish plaques on pressure areas of the soles (b–d) and a defining feature of the condition, usually develops after formation of blisters. The reaction appears to be dose-dependent. The photographs show increasing degrees of severity from grade I (a) to grade II (b) and grade III (c and d) (reproduced with permission from Degen et al. [151]).
cyclophosphamide [179], pentostatin [44] and suramin [180] have already been shown to induce immune thrombocytopenia. Evan’s syndrome (acute thrombocytopenia with haemolysis) to oxaliplatin has also been described [181, 182]. Early onset following previous drug exposure and the detection of antibodies and complement on erythrocytes indicated a type II hypersensitivity reaction [182].

An interesting new mechanism involving splenic sequestration of platelets during oxaliplatin-induced liver damage has recently been identified in some cases of oxaliplatin-related thrombocytopenia [183]. The capacities of the older anti-neoplastic drugs to induce dose-dependent thrombocytopenia are generally well-known—those that do not lead to platelet reduction or do so only weakly include L-asparaginase, bleomycin, cisplatin, flutamide, goserelin, luprolide, megesterol, streptozocin, tamoxifen and vincristine; drugs causing moderate thrombocytopenia include actinomycin D, cyclophosphamide, 5-fluorouracil, 6-mercaptopurine, methotrexate and procarbazine and drugs that induce the condition in severe form include busulfan, chlorambucil, cytotoxic arabinoside, melphalan, nitrogen mustard, nitrosoarene compounds and vinblastine. The propensities and relative potencies of many of the more recently introduced anti-neoplastic drugs (especially many used in targeted therapies such as signal transduction and proteasome inhibitors) to cause both non-immune and immune thrombocytopenia do not appear to have been widely studied as yet, and little comparative data are available. Clearly this is important information to know since while dose-dependent non-immune thrombocytopenia can often be controlled or avoided by a reduction in dose, drugs causing immune-mediated thrombocytopenia must be avoided.

5.2.2 Chemotherapy-induced neutropenia

Drug-induced neutropenia (DIN) is probably the most frequent cause of neutropenia. Neutropenia is the most common adverse response induced by anti-cancer drugs, and neutropenia with fever is a major dose-limiting effect [184] of many of these drugs. Febrile neutropenia is generally defined as a rise in axillary temperature to more than 38.5 °C for more than 1 h. These drugs are essentially all anti-neoplastic drugs well-known for their association with neutropenia (especially many used in targeted therapies, such as signal transduction and proteasome inhibitors) to cause both non-immune and immune thrombocytopenia do not appear to have been widely studied as yet, and little comparative data are available. Clearly this is important information to know since while dose-dependent non-immune thrombocytopenia can often be controlled or avoided by a reduction in dose, drugs causing immune-mediated thrombocytopenia must be avoided.

5.2.3 Drug-induced haemolytic anaemia

The well-known immune cytotoxic reaction to penicillin resulting from binding of the drug to red cells, reaction of IgM and IgG antibodies with the drug–cell membrane protein complex and activation of the classical complement pathway causing cell lysis and death is the prototype drug-induced type II hypersensitivity. Another example of a type II cytotoxic antibody-mediated drug reaction, commonly seen with some cephalosporins, is drug-induced immune haemolytic anaemia (DIIHA) where the drug appears to form an antigenic complex...
with the red cell surface [189]. In the analysis of 75,243 chemotherapy regimens of associated anaemia and thrombocytopenia mentioned above, from 46.4 to 59 % of treated patients developed anaemia [173]. DIIHA can also be associated with red cell autoantibodies without the drug participating in the antigen–antibody reaction. The anti-neoplastic drug fludarabine is an example of such a drug [43, 190, 191], and other purine analogues, cladribine [41, 192] and pentostatin [46], have also caused DIIHA. The closely related platinum drugs cisplatin, carboplatin and oxaliplatin are known to cause immune-mediated haemolysis with or without associated immune thrombocytopenia [193–196]. It is not fully understood why some drugs sometimes induce antibodies to red cells or what mechanism(s) is involved [189]. A possible immune basis for any anti-neoplastic drug that induces anaemia should always be kept in mind. The autoimmune disease bullous pemphigoid [197], classified as a type II hypersensitivity, can be caused by a number of drugs including the selective oestrogen receptor modulator (SERM) tamoxifen.

5.3 Type III reactions

These hypersensitivity reactions to chemotherapeutic drugs involving immune complex formation occur less often than type I, II or IV reactions, but it seems they also go unrecognised more often.

5.3.1 Drug-induced vasculitis

Drug-induced vasculitis (DIV) usually occurs in the skin and sometimes in subcutaneous tissue, kidneys and the lungs. The most common clinical symptoms at onset include skin rash, arthralgia and myalgia. For anti-neoplastic drugs, small vessel vasculitis is the most frequently seen form of a type III reaction. Cutaneous vasculitis (CV) is a small vessel systemic vasculitis manifesting as palpable purpura and ranging in severity from being benign and self-limiting to life-threatening with multiple organ failure. CV seen in malignancy may be drug- or infection-induced or paraneoplastic in origin. Approximately 60 % of patients with DIV present as CV [198]. Mechanisms underlying DIV are still incompletely understood, but cellular as well as humoral immune processes appear to be involved. A proportion of small-vessel vasculitis patients have anti-neutrophil cytoplasmic antibodies (ANCA) which are used as a diagnostic marker. In ANCA-positive DIV patients where neutrophil damage may occur, antibodies to myeloperoxidase, cathepsin G, human leukocyte elastase and lactoferrin may be found [199]. These antibodies can be indicative of severe disease. Because of the presence of the many different autoantibodies and low complement C4 levels, DIV may be confused with lupus [200]. Chemotherapy drugs implicated in CV include aromatase inhibitors (e.g. anastrozole) [115], SERMs such as tamoxifen [200, 201], EGFR inhibitors including erlotinib which has been involved in at least 14 cases [202] and cyclophosphamide [203].

Paraneoplastic vasculitis, an inflammatory reaction of the vessels induced by malignant cells [204], represents about 5 % of cases of CV. Ninety percent of cases occur in haematological malignancies. The reactions appear to be the result of paraproteinemia (usually cryoglobulins), but the pathogenesis is unclear. One suggestion is that immune complexes form with tumour antigens leading to release of lymphokines and other vasoactive substances that damage vascular endothelium [205].

5.3.2 Other drug-induced type III reactions

Oxaliplatin has been associated with type III immune complex-mediated urticaria, joint pain and proteinuria [206] and idiosyncratic reactions including pulmonary fibrosis and cytokine release syndrome where it is speculated that it may act as a superantigen [207]. A reaction consisting of fever, myalgia, bone pain, conjunctivitis, chest pain and maculopapular rash induced by cytarabine in patients with NHL or ALL and termed cytarabine syndrome [208] has been suggested to be a type III hypersensitivity response on the basis of the detection of immune complexes and successful treatment with corticosteroids. Procarbazine has been said to induce a type III reaction with immune complex deposition ‘manifesting as a toxic epidermal necrolysis’ [209], a reaction normally considered to be a type IV response. A possible type III hypersensitivity reaction was also the tentative conclusion in each case from studies of procarbazine-induced interstitial pneumonia with eosinophilia improved by corticosteroid administration, viz. patients with diffuse pulmonary infiltrates, alveolar damage and eosinophilia after receiving gemicitabine; haemolytic anaemia and a Henoch–Schönlein-type purpura each associated with immune complexes following mitomycin C administration; and pneumonitis with features of a type III hypersensitivity reaction was also the tentative conclusion in each case from studies of procarbazine-induced interstitial pneumonia with eosinophilia improved by corticosteroid administration, viz. patients with diffuse pulmonary infiltrates, alveolar damage and eosinophilia after receiving gemicitabine; haemolytic anaemia and a Henoch–Schönlein-type purpura each associated with immune complexes following mitomycin C administration; and pneumonitis with features of a type III response provoked by methotrexate [21]. A further discussion of possible lung hypersensitivities to chemotherapeutic drugs is set out below for drug-induced lung disease (DILD). An unusual example of a possible type III hypersensitivity reaction to a chemotherapeutic drug was demonstrated by bronchial challenge testing an asthmatic nurse occupationally exposed to mitoxantrone in an oncology ward. The patient responded to the drug with a biphasic reaction and an increase in eosinophils, neutrophils and lymphocytes 6 to 18 h post-challenge [21].

5.4 Type IV reactions

Less often seen cutaneous reactions to chemotherapeutic drugs include maculopapular exanthemas, allergic contact dermatitis, psoriasis, acute generalized exanthematous pustulosis (AGEP), drug reaction (rash) with eosinophilia and systemic
symptoms (DRESS; also termed drug-induced hypersensitivity syndrome), fixed drug eruptions, erythema multiforme, Stevens–Johnson syndrome (SJS) and toxic epidermal necrolysis (TEN). These are now generally considered to be immune-based, delayed type IV hypersensitivities mediated by antigen-specific effector T cells. Type IV reactions generally begin from about 7 to 21 days after contact with the drug, and subsequent reactions may appear only 1 or 2 days after re-exposure. Identification and specificity of the culprit drug is established by oral challenge, patch and intradermal tests (IDT) read after a delay of at least 48 h [170]. Investigations are proceeding to relate different T cell subsets and cytokine and chemokine profiles with different skin reactions although cytokine overlap occurs [210]. There are many more examples of type IV cutaneous reactions to non-targeted than to targeted anti-neoplastic drugs although this might reflect, at least in part, the more recent introduction and so far smaller number of recipients of the latter drugs. The main drugs involved in eliciting delayed skin reactions include alkylating agents, particularly the nitrogen mustards; some anti-metabolites, e.g. cytarabine, gemcitabine, some purines; the folate antagonist methotrexate and the taxanes. Tamoxifen, used for many years, has been implicated in erythema multiforme and SJS. Other targeted drugs so far implicated in the induction of type IV skin reactions include imatinib and vandetanib in SJS, bortezomib, imatinib, letrozole and pralatrexate in TEN, exemestane, imatinib and letrozole in erythema multiforme, exemestane and imatinib in AGEP, imatinib in DRESS and bexarotene, erlotinib and sorafenib in reactions defined as ‘bulous’ eruptions (Table 3).

5.5 Other drug-induced reactions showing some evidence of hypersensitivity responses

Some adverse reactions provoked by chemotherapeutic drugs, for example, in the skin, liver, vasculature and lungs, show features of antibody and/or cellular involvement that suggest a possible contribution of type II, III and/or IV hypersensitivity mechanisms but, for a number of reasons, clear and definitive indications to establish such classifications one way or the other are often lacking. In addition to the lack of research so far, at least part of the problem in defining some of these adverse reactions is due to the lack of a generally accepted definition of hypersensitivity, the frequent absence of tests to identify or discount allergic recognition and the often vague classifications of reactions without any apparent consideration of the possible involvement of types II, III and IV hypersensitivity responses. These factors are discussed further below. Some of the more important drug-induced reactions that show at least some features of true hypersensitivity reactions but which are not routinely included in the Gell and Coombs classification of hypersensitivities will now be considered from both immune and allergological perspectives.

5.5.1 Some drug-induced cutaneous reactions

As already discussed, cutaneous reactions to anti-neoplastic drugs are common and may range from mild rashes and urticaria to severe type IV cell-mediated toxidermias such as erythema multiforme, SJS and TEN. Examples of antibody-mediated cutaneous reactions are seen in drug-induced cutaneous vasculitis, a type III hypersensitivity reaction, and some cases of urticaria are true type I IgE antibody-mediated hypersensitivities. Other cutaneous reactions where a hypersensitivity mechanism is less obvious include site-specific toxicities involving the hair, nails, mouth, hands and feet all of which are commonly seen. While the aim of chemotherapy is to kill rapidly growing cancer cells, normal rapidly growing cells in the skin, nails and hair follicles are also inhibited. It is not surprising therefore that adverse reactions such as alopecia, simple macular rash, xerosis cutis, pruritus, hand–foot skin reaction, pigmentary changes and nail dystrophies occur often in patients undergoing cancer chemotherapy.

Apart from the well-known types I, II, III and IV hypersensitivity reactions, major cutaneous and mucocutaneous reactions associated with chemotherapy include stomatitis and mucositis, acral erythema, alopecia, cutaneous eruption of lymphocyte recovery, eccrine squamous syringometaplasia (ESS), extravasation reactions (including chemical cellulitis), hyperpigmentation, inflammation of keratosis, nail reactions, neutrophilic eccrine hidradenitis (NEH), photosensitivity and radiation associated (recall and enhancement) reactions [209, 211]. Mechanisms underlying most of these reactions remain to be determined. Cytotoxic chemotherapeutic drugs are the most common cause of stomatitis and mucositis. Although these terms are often used synonymously, stomatitis (oral mucositis) is restricted to the oral mucous membranes while mucositis affects the GI tract from mouth to anus. Approximately 40 % of patients receiving chemotherapy experience stomatitis. The most stomatotoxic agents include alkylating agents busulfan, chlorambucil and cyclophosphamide, anti-metabolites cytarabine, 5-fluorouracil and methotrexate, antibiotics bleomycin, actinomycin D, doxorubicin and mitomycin C, Vinca alkaloids and the taxanes [212]. Cytotoxic anti-neoplastic drugs destroy the rapidly dividing cells of the oral mucosa at a faster rate than most other normal healthy cells, but the detailed pathogenetic mechanism of stomatitis is still to be worked out. The pathogenesis of acral erythema (palmar–plantar erythrodysesthesia; hand–foot syndrome) [153] is not known, but it may be a direct cytotoxic effect. Some suggested mechanisms to explain the palms–soles location (Figs. 3 and 4) of reactions are based on vascular anatomy, temperature gradients, rapidly dividing epidermal cells and high concentration of eccrine glands [209]. Cutaneous eruptions of lymphocyte recovery is seen as a sharp, transient rise in temperature accompanied by macular and papular eruptions during the earliest recovery in peripheral lymphocyte numbers.

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following chemotherapy for leukemia [213]. The observed rash is thought to indicate the return of immunocompetent lymphocytes to the circulation suggesting an immune pathogenesis. ESS is a benign cutaneous reaction not associated with any particular cytotoxic chemotherapeutic drug which manifests as erythematous macules, plaques or vesicles on the intertriginous areas of the palms and soles. The reaction is thought to be a direct cytotoxic effect of the drugs. Extravasation reactions occur when chemotherapeutic drugs are inadvertently allowed to contact the tissues during their IV administration. Reactions may be irritant or vesicant, the former appearing as phlebitis and an erythematous reaction along the vein or at the IV site; urticarial flare reactions exhibited by anthracyclines are an example. A well-known drug eliciting vesicant reactions (called chemical cellulitis) is the antibiotic mitomycin C. Several drugs, including suramin, cytarabine and 5-fluorouracil, provoke inflammation in pre-existing skin reactions such as actinic keratoses. A suggested mechanism is increased DNA synthesis and uptake of drug by the affected tissues [214]. NEH [215], associated with cytarabine, bleomycin, chlorambucil, cyclophosphamide, mitoxantrone, vincristine and a number of other cytotoxic anti-cancer drugs, appears as fever, oedema and skin eruptions such as erythematous macules, papules, plaques and pustules. Necrosis of eccrine epithelial cells together with neutrophils around eccrine sweat glands and ducts and the presence of drug in the sweat of patients suggest that direct toxicity of the sweat glands is the cause of NEH [216].

Although modern targeted anti-cancer therapies such as tyrosine kinase and proteasome inhibitors have seen reductions in common systemic toxicities (e.g. nausea, vomiting, asthenia, diarrhoea etc.) and some of the more severe cutaneous and gastrointestinal effects occurring with some myelosuppressant drugs, it has become apparent that the targeted agents are responsible for multiple mild to moderate dermatological adverse effects (Table 3) [152, 155]. The most commonly reported effects are hand–foot skin reaction [151] (Fig. 4) (a distinct entity from hand–foot syndrome or acral erythema seen with cytotoxic drugs such as capecitabine (Fig. 3), 5-fluorouracil and doxorubicin [152]), papulopustular rash (Fig. 5), xerosis, fissures on fingertips and toes, pruritus, pigmented changes and abnormalities of the mouth (stomatitis, mucositis), hair, scalp (alopecia), nails (e.g. paronychia) and on the eyelids (trichomegaly of the lashes). Some of the main features that can be used to distinguish the two hand–foot reactions relate to the type of lesion and their location on the hands and feet. The cytotoxic drugs cause diffuse and symmetric erythematous lesions with oedema and fissuring while in hand–foot skin reactions, tender, localized lesions appear as blisters and hyperkeratosis [152]. Lesions are seen on the palmar (ventral) surfaces of the hands following administration of both types of drugs. They are diffuse in the case of the cytotoxic agents but occur on the digit tips, over the interphalangeal joints and on the thenar area following the expression of other cutaneous reactions. In hand–foot skin reaction [151] (Fig. 4).
DILI is predominantly seen as hepatitis and/or cholestasis [217]. Most cases of DILI fall into three main clinical patterns: acute hepatocellular injury with jaundice similar to hepatitis and increased serum transaminase; acute cholestatic injury with jaundice, marked elevation of serum alkaline phosphatase and moderate increases in transaminases and a mixture of these two patterns showing elevations of both alkaline phosphatase and transaminases [217]. The mechanism of DILI can be divided into direct, sometimes called intrinsic, hepatotoxicity (as seen with, for example, paracetamol (acetaminophen)) and unpredictable and rare liver injury which in turn can be classified into immune-mediated hypersensitivity and idiosyncratic reactions [217, 219]. The latter reactions are only seen in susceptible individuals and may be due to idiosyncratic metabolism producing harmful metabolites. Allergic hepatitis is associated with fever and rash and when caused by drugs, as in for example, DRESS, the reaction is generally a type IV hypersensitivity response involving CD4+ cells, CD8+ cytotoxic lymphocytes and NK, Kupffer and dendritic cells. Type II hypersensitivities may also sometimes occur. There are currently two main hypotheses for the mechanism of immune-mediated DILI. In perhaps the most supported model, the drug or active metabolite(s) as hapten is said to bind to endogenous proteins forming immunogenic conjugates that generate antibody- and/or T cell-mediated injury. Another view is that most individuals are tolerant to immune injury of the liver and reactions occur only when tolerance is in some way overcome [220–222]. Knowledge of mechanisms underlying idiosyncratic DILI remains limited. Direct hepatotoxicity usually manifests as necrotic reactions with little inflammation while idiosyncratic reactions show injury with clear inflammation. Anti-neoplastic drugs associated with an acute hepatitis injury pattern include L-asparaginase, azathio- prine, carmustine, cyclophosphamide, 6-mercaptopurine, mithramycin and vincristine.

In considering DILI based on histological patterns and differential diagnoses, Ramachandran and Kakar [219] have distinguished and classified the involvement of a number of different commonly used chemotherapeutic drugs. In chronic hepatitis lacking autoimmune markers, 5-fluorouracil, its produg tegafur and tamoxifen have been implicated. Progression to fibrosing cholestatic hepatitis was reported in a hepatitis C patient given cyclophosphamide and corticosteroids [223]. In autoimmune hepatitis accompanied by hypersensitivity, rash, eosinophilia and arthralgia, long-term use of the tetracycline minocycline may lead to autoimmune hepatitis or hepatitis that mimics lupus-related hepatitis. The risk of methotrexate-induced liver toxicity is exacerbated by high dosage, existing liver disease and heavy alcohol use. The alkylating agent procarbazine, metabolised and activated in the liver, may cause granulomatous hepatitis. Methotrexate, 5-fluorouracil, cisplatin and tamoxifen have been associated with macrovesicular steatosis while the topoisomerase I inhibitor irinotecan has a direct effect in steatohepatitis. Such an effect, sometimes referred to as chemotherapy-associated steatohepatitis, has also been reported for oxaliplatin. Tamoxifen, oestrogen and diethylstilbestrol can lead to hepatic steatosis but rarely steatohepatitis. Oxaliplatin may injure sinusoidal endothelial and hepatic stellate cells; thiopurine chemotherapeutic drugs have been implicated in peliosis hepatis and dacarbazine in hepatic vein thrombosis presenting clinically as Budd–Chiari syndrome [219].

5.5.3 Drug-induced lung disease

The term DILD covers a heterogeneous group of lung diseases. Patients most likely to develop DILD are those receiving chemotherapy with up to about 10% of such patients experiencing injury although the incidence for individual drugs is different. The anti-neoplastic drugs most commonly involved in DILD are bleomycin, methotrexate, cyclophosphamide and busulfan, but many chemically and mechanistically different anti-cancer agents demonstrate pulmonary toxicity including mitomycin C, chlorambucil, melphanal, 6-mercaptopurine, cytarabine, gemcitabine, fludarabine, carmustine, lomustine, etoposide, the taxanes, irinotecan, procarbazine, vinblastine, zinostatin and the tyrosine kinase inhibitors imatinib and gefitinib. In general, mechanisms involved in the drug-induced lung injuries have not been elucidated making it difficult to effect classifications on the basis of pathogenesis. Chemotherapeutic drugs can be grouped into four main divisions: those producing interstitial pneumonia and fibrosis, hypersensitivity reactions, acute respiratory distress syndrome (ARDS) and bronchiolitis obliterans organizing pneumonia (BOOP) [224]. Interstitial pneumonia and fibrosis, often provoked by bleomycin, is an inflammation of the lung interstitium and the most commonly occurring DILD. Other drugs involved include busulfan, chlorambucil, cyclophosphamide, melphanal, methotrexate and paclitaxel [225]. DILD hypersensitivity reactions result from interaction of drug with the immune system. Signs of interstitial pneumonia with lymphocytes and plasma cells infiltrating the interstitial space are seen. Immune-mediated damage to the lung in DILD may be due to drug-specific antibodies or, more usually, drug-specific T cells. Eosinophilic pneumonia showing eosinophils in the peripheral blood and bronchoalveolar lavage (BAL) fluid can be caused by almost any medication. Hypersensitivity pneumonia is a combined types III and IV reaction in a Th1/Th17 response [226]. Reports of drug-induced hypersensitivity pneumonitis are increasing particularly to anti-neoplastic drugs [226]. No single diagnostic test is sufficient for a diagnosis, rather a combination of careful history, imaging (e.g. high-resolution computed tomography (CT), examination of BAL fluid (positive signs include lymphocytic alveolitis with an increase in CD8+ lymphocytes), serum
drug-specific antibody tests, lung biopsy and perhaps inhalation challenge and the lymphocyte transformation test (LTT). Corticosteroids appear to hasten resolution of symptoms in both drug-induced hypersensitivity and eosinophilic pneumonia. In ARDS, inflammation of the lungs leads to impaired gas exchange and release of mediators causing further inflammation, hypoxia and ultimately organ failure. All trans-retinoic acid (ATRA) and mitomycin C are examples of chemotherapeutic drugs causing ARDS. In BOOP, organized polyloid granulation inflammatory tissue is seen in the distal bronchi-ole Airways, respiratory bronchioles, alveoli and alveolar ducts [227, 228]. Bleomycin, cyclophosphamide and methotrexate are common causes.

The lack of definitive diagnostic tests for suspected chemotherapy-associated hypersensitivities makes it difficult to definitively identify some lung reactions thought to have an immune or hypersensitivity basis (see below), but some reactions appear to be true hypersensitivity responses. One such example is an acute onset form of bleomycin pneumonitis with fever and peripheral blood and BAL eosinophilia reversed by corticosteroids. Other examples include methotrexate-associated pneumonitis, an acute hypersensitivity reaction with bronchospasm to gemcitabine, procarbazine pneumonitis and pulmonary infiltrates and interstitial pneumonitis following docetaxel and paclitaxel [229, 230]. Some mitomycin C-treated adenocarcinoma patients with an unusual haemolytic–uraemic-like syndrome and pulmonary oedema develop ARDS. Circulating immune complexes with antibodies to carcinoembryonic antigen, platelets and gastric carcinoma cell surface antigens detected in a few of the patients suggest that the syndrome may be initiated by tumour cells or cell products released as a result of chemotherapy [231–233].

6 Cytokine-release and tumour lysis syndromes

Cytokine-release syndrome (CRS), sometimes called acute infusion reaction, is different from a hypersensitivity reaction, but the two may share some signs and symptoms, for example, nausea, fever, cough, dyspnea, bronchospasm, hypotension, rash, itching and urticaria. CRS usually develops during or soon after drug infusion with symptoms resolving within about 24 h [234]. Destruction of target and immune effector cells is thought to release cytokines such as TNF and interleukins into the circulation producing symptoms similar to a type I hypersensitivity response. Oxaliplatin, for example, has been implicated in the release of TNF-α and IL-6 in five colorectal cancer patients experiencing the signs and symptoms of CRS after infusion of the drug [235]. Symptom regression in the patients paralleled the decrease in serum cytokine concentrations. Prompt interruption of infusion and/or therapy with antihistamines, IV fluids, NSAIDs and narcotics is required for CRS, and it should be continued prophylactically for at least 24 h [234].

Tumour lysis syndrome (TLS) occurs 48–72 h after initiation of cancer therapy when large numbers of tumour cells are killed in a short time, releasing intracellular contents into the circulation. This rapidly results in an ionic imbalance involving hyperkalaemia, hyperphosphatemia, hypocalcaemia and hyperuricaemia which may progress to acute renal failure, cardiac arrhythmias, seizures and death [236]. TLS occurs most often in ALL and lymphoma patients but is also seen with solid cancers particularly small-cell cancer and neuroblastoma. Anti-neoplastic drugs known to cause TLS include etoposide, fludarabine, paclitaxel, hydroxyurea, zoledronic acid, thalidomide and bortezomib. Unlike CRS, TLS is not difficult to differentiate from drug-induced hypersensitivity responses.

7 Diagnostic approaches and methods

7.1 Skin testing with chemotherapeutic drugs

The skin prick test (SPT) and IDT may be used to detect both immediate and delayed reactions to drugs, but these tests have not been widely and routinely applied, being used to diagnose hypersensitivities to only a relatively few drugs including platin, taxanes, cyclophosphamide, ifosfamide, cytarabine, 5-fluorouracil, methotrexate, mitomycin C and epirubicin [18, 19, 21]. Given the many non-targeted anti-cancer agents, let alone the ever-expanding range and variety of new targeted drugs, it is obvious that skin testing remains significantly underinvestigated and under utilized for diagnosis of drug hypersensitivities in oncology. Patch testing is said to be rarely, if ever, positive and not considered useful [24] (see, however, section 7.2.7).

Skin testing has been employed more often with the platin than any other anti-neoplastic agent but, even with these drugs, the procedure has not been widely used as a routine diagnostic test. ID testing with 100–240 μg of carboplatin showed that while only 13 of 47 patients (28 %) with ovarian or peritoneal cancer had a positive reaction, a negative skin test correctly predicted the absence of hypersensitivity in 166 of 168 courses of chemotherapy (658 of 668 patients or 98.5 %) [237]. A follow-up study confirmed the association between a negative carboplatin skin test and absence of hypersensitivity, but the significance of a positive test remained less certain [238]. Positive test results were obtained more often in a group of 60 patients referred for reactions to carboplatin. Fifty-three patients were skin test-positive, one patient showed a delayed positive reaction, four responded with a hypersensitivity reaction during drug infusion and two proved positive after several more infusions [166]. In a more recent study, positive skin tests to carboplatin were obtained in 64 % of 58 patients, leading the authors to conclude that the
test is not a reliable predictor of carboplatin hypersensitivity [239]. Skin test concentrations generally employed for carboplatin are 10 mg/ml in the SPT and 0.02 ml of 0.1 or 1 mg/ml in the IDT increasing in ten-fold concentration steps to a maximum of 10 mg/ml. For oxaliplatin, concentrations of 1–5 mg/ml have been used in the SPT and maximums of 0.1–5 mg/ml in the intradermal test [165, 240]. In summary, it has been claimed that skin tests are positive in more than 80 % of the platinum drug-treated patients tested, and when the skin test to platinum drugs is negative, the risk of a hypersensitivity reaction is reduced sevenfold or even eliminated [57]. This has led to the recommendation that skin testing should be performed on every patient before the eighth drug infusion.

For the diagnosis of hypersensitivity to oxaliplatin, skin testing is said to be positive in 75 to 100 % of patients [164, 165]. Skin testing may also help in ruling out possible cross-reactions when seeking to substitute one platinum drug for another [241]. Employment of skin testing in everyday oncology practice remains controversial, principally because of lack of prior experience, the need for controls subjects and the fact that reactions may still occur in patients with a negative test [57]. This draws attention to the need for some allergy training in clinicians administering chemotherapy (see below).

7.2 Testing for types I, II, III and IV hypersensitivities

In general, diagnosis of type I hypersensitivities to chemotherapeutic drugs is based on clinical assessment and only sometimes with the aid of skin testing and rarely challenge testing. Specific assays for the detection of chemotherapeutic drug-specific IgE antibodies are usually not available, and when occasionally used, they are generally not validated. The basophil activation test is yet to be systematically investigated and applied for mechanistic studies or diagnosis.

7.2.1 Drug-induced thrombocytopenia

As mentioned above in relation to drug-induced thrombocytopenia, anaemia and neutropenia, it is important to distinguish immune-based suppression of haematopoietic cell lines from the dose-dependent bone marrow cytotoxicities shown by many anti-neoplastic drugs, but this is not always easily done. Since thrombocytopenia in response to chemotherapy is commonly the result of marrow suppression of megakaryocytopenia, a well-recognized side effect of anti-neoplastic drugs, immune-mediated DIT is not always considered in the diagnosis. Diagnosis, even harder if the patient is taking multiple medications, often tends to be based on the temporal relationship between drug administration and symptoms and/or on one-at-a-time drug withdrawal along with examination for a possible rebound in the platelet count. In vitro tests in the form of immunoassays that detect platelet-reactive serum antibodies are available [242, 243], but while they are an aid in the diagnosis of idiopathic thrombocytopenic purpura and autoimmune DIT, the tests are not standardized, often present technical difficulties (such as poor aqueous solubility of the drug and questions of the involvement of metabolites) and do not generally detect the drug-dependent antibodies. Specific tests in the form of enzyme-linked immunoassays (ELISAs) and cytometric procedures for drug-dependent antibodies in individual patients’ sera are available at a few reference laboratories, but limited access to them and the time taken to obtain the result in an urgent situation are major drawbacks. Also, in some cases, a false negative result may result if the patient’s own platelets are not employed in the test. Recommendations for the implementation of platelet autoantibody testing for immune thrombocytopenia have been published [243].

7.2.2 Drug-induced neutropenia and agranulocytosis

In addition to a complete blood count and differential and manual examinations of peripheral blood smears, anti-neutrophil antibody tests may be utilized to help diagnose immune-mediated neutropenia and agranulocytosis. Anti-neutrophil antibodies to neutrophil glycoproteins are detected by several different immunoassays [186]. The neutrophil antigens involved in immune neutropenia have been given the nomenclature HNA-1a, HNA-1b and HNA-1c (glycoprotein FcyIIIb; CD16), HNA-2a (gp50–64; CD177), HNA-3a (gp70–95), HNA-4a (CD11a) and HNA-5a (CD11b). Immunoassays employed include a granulocyte agglutination test, direct and indirect granulocyte immunofluorescence tests, ELISAs and, perhaps the most specific of the tests, the monoclonal antibody immobilization of granulocyte antigens (MAIGA) assay [244]. Unfortunately, tests for neutrophil antibodies are not widely performed and available, there are a number of technical difficulties including the detection and distinguishing of autoantibodies and drug-dependent antibodies [187]. Fc receptors on neutrophils can produce false positive results, cells show fragility, aggregate and lyse and test results are not always easy to interpret. Taken together, these factors help to explain why ready detection of antibodies to neutrophils and hence easy diagnosis of cases of immune-mediated drug-induced neutropenia/agranulocytosis is not necessarily easy and sometimes not achievable.

7.2.3 Drug-induced anaemia

Tests for drug-induced anaemias are not as problematic as those for platelets and neutrophils, but the situation can be more complicated than it appears at first sight. Haemolytic anaemia (HA) resulting from high doses of penicillin is the classical cytotoxic type II hypersensitivity. Binding of drug to erythrocytes causes the cells to be recognized as foreign resulting in IgM and IgG antibodies reacting with the drug-
cell membrane protein complex, activation of the classical complement pathway, cell lysis and death. DIIHA [189] is another example of a type II cytotoxic antibody-mediated drug reaction. It can be associated with red cell autoantibodies that are drug-independent, that is, detected in vitro without adding drug or with drug-dependent antibodies that only react in vitro in the presence of drug. Some drugs bind covalently to the erythrocyte membrane proteins, antibodies bind to the surface-bound drug and following interaction with macrophages, Fc-mediated destruction of red cells occurs. The mechanism with other drugs remains controversial. Fludarabine is the most common drug causing drug-independent red cell autoantibodies, but other anti-neoplastic purine analogues, cladribine and pentostatin and the platinum-based chemotherapeutics have also caused DIIHA. Drugs already in use or newly introduced and not yet known to cause anaemia will be implicated in the future. Detailed mechanistic studies of most cases of anti-neoplastic drug-induced HA are generally not routinely undertaken so it is possible that interesting and surprising findings may be revealed with some of the many drugs now employed, as the following example serves to illustrate. In a study of 300 CLL patients given fludarabine, cyclophosphamide and rituximab, 5.8 % developed HA but 82.4 % of these patients showed a negative direct antiglobulin test suggesting that the drug combination treatment led to antiglobulin test-negative autoimmune HA [245].

7.2.4 Drug-induced vasculitis

Diagnosis of DIV can be difficult since there are no clearly well-established clinical or laboratory markers that can distinguish the condition from other vasculitidis. Anaemia is common in patients with DIV, pulmonary symptoms may correlate poorly with the disease and CT scanning of the chest, an ANCA screen (see section 5.3.1) and tissue biopsy may be necessary to reach a definitive diagnosis [246]. Some patients may also have antibodies to phospholipids and/or histones.

7.2.5 Drug-induced liver injury

As mentioned earlier, most cases of DILI fall into three main patterns [217] which may be used along with historical patterns [219] for differential diagnoses. Individual drugs often tend to have their own pattern signature that is characteristic of the drug [217]. Injury features such as rash, eosinophilia and a rapid response on re-challenge suggest that so-called idiosyncratic cholestatic liver injury is allergic in nature [217]. Accurate diagnosis of DILI, especially immune-mediated DILI together with identification of the causative drug, remains difficult. Specific tests are not available and there are no relevant experimental models to confidently establish mechanisms and devise potential diagnostic procedures. Current assessment of DILI is therefore largely dependent on serum concentration of total bilirubin and tests for liver damage namely, measurements of serum alkaline phosphatase and liver transaminases alanine transaminase and aspartate transaminase. For immunological tests, LTT has been used for the detection of T cell proliferation, but the test is not always reliable and not always regarded with confidence if only because of doubts about antigen presentation of different drugs [221].

7.2.6 Drug-induced lung disease

Diagnosis of immune-mediated DILD provoked by chemotherapy drugs is similar to the situation with DILI. Although both an in vivo and in vitro approach would seem to be applicable, neither is totally satisfactory in practice. As discussed earlier, chemotherapeutic drugs can elicit four main types of lung reactions. High-resolution CT scanning, pulmonary function testing and bronchoscopy with BAL can be used along with the patient’s history of drug exposure, histological evidence of lung damage and exclusion of other causes [224, 227]. In CT scans, interstitial pneumonitis and fibrosis (e.g. caused by bleomycin) tend to involve the lower lung zones, hypersensitivity reactions may be seen as ground-glass opacities with centrilobular nodules, bilateral ground-glass opacities involving dependent lung regions may be seen in ARDS and peribronchial or subpleural areas of consolidation are seen in BOOP [224]. In vitro, the LTT has been quite widely used in the attempted diagnosis of DILD in Japan where it has been said that ‘compelling data as to the sensitivity and specificity of the (test) is currently lacking’ [247]. For methotrexate, the LTT has been shown to be inadequate for confirming DILD. Positive results have been claimed for a leukocyte migration test designed to identify cytokines or chemokines produced by drug-stimulated lymphocytes from patients with methotrexate- or paclitaxel-induced pneumonitis. However, the sample numbers are so far small [247]. In vivo drug provocation testing is generally considered too risky since pulmonary damage may be irreversible and the utility of skin testing with drugs in DILD does not appear to have been carefully and systematically investigated.

7.2.7 Cutaneous reactions

As discussed, anti-cancer drug therapies are associated with multiple and various dermatological effects some of which are clearly immune-mediated, usually type IV hypersensitivities, and others which are less well understood mechanistically, especially those associated with targeted therapies such as EGFR inhibitors. With the latter drugs, cutaneous eruptions such as hand–foot skin reaction, papulopustular rash, lichenoid drug eruptions and palmoplantar hyperkeratosis [86, 152, 248] are in need of investigations to identify underlying mechanisms and to establish whether or not immune
processes have any role at all in drug-induced dry skin, pruritus and mouth, scalp, hair and nail changes.

Patch testing with drugs, in both pure and commercial form, is used to determine the cause of drug-induced cutaneous drug reactions and for studying the underlying pathophysiological mechanisms. The test is both a screening test for hypersensitivity and a provocation test in the target organ [170], the skin, where it can be seen as reproducing the disease. Patch testing is valuable for investigating a number of skin reactions including eczema, contact dermatitis, maculopapular rash, photosensitivity, fixed drug eruption, lichenoid rash and acute generalized exanthematous pustulosis. It seems to be less useful for investigating urticaria, SJS and TEN. Patch testing with anti-neoplastic drugs is generally considered to be of no value in diagnosing systemic reactions (see section 7.1), and it remains unclear how useful the test is for helping in the diagnosis of chemotherapeutic drug-induced cutaneous reactions. The LTT measures a memory T cell response and may identify a causative drug in cases of drug eruptions, but it is important to perform the test at the right time. The time varies and depends on the reaction—for example, with DRESS, patients should be tested 5 to 8 weeks after the onset of reactions while for maculopapular drug eruptions, SJS and TEN, testing should take place within 1 week of skin rashes. Other promising in vitro approaches for diagnosing and studying immune-mediated, in particular type IV, cutaneous reactions to drugs include application of the local lymph node assay for identifying contact allergens, ELISPOT cytokine assays for the detection of drug-reactive T cells and monitoring cutaneous lymphocyte-associated antigen and levels of the skin-associated chemokine CCL27 and its interaction with its receptor CCR10 [170].

8 Premedication and desensitization for chemotherapy

There is a risk of hypersensitivity reactions following administration of some drugs used in chemotherapy, and to minimize the risk and reactions, premedication may be undertaken. Taxane therapy, for example, requires premedication but premedication does not prevent true type I reactions to the platins drugs [24]. Reactions to taxanes may be IgE antibody-mediated or due to direct mast cell/basophil or complement activation. Corticosteroids and antihistamines are generally employed in premedication protocols. Patients given paclitaxel require IV H1 (diphenhydramine 50 mg) and H2 (cimetidine 300 mg, ranitidine 50 mg, famotidine 20 mg) antagonists prior to infusion commencing and an oral steroid (dexamethasone 20 mg) 12 and 6 h prior to 1-, 3- and 24-h-long infusions. For patients on docetaxel, dexamethasone 8 mg is given twice daily for 3 days starting 24 h prior to the commencement of infusion [249]. Premedication does not seem to be required for infusions extending over 96 h or more. In one protocol for docetaxel given weekly, oral dexamethasone 4.5–7.5 mg is administered twice daily on the day before, the day of and the day after commencement of infusion, together with IM promethazine 25 mg and IV cimetidine 600 mg 30 min before docetaxel. For paclitaxel, the protocol is: dexamethasone 2.25–7.5 mg orally 12 h and 2 h before and promethazine 25 mg IM and cimetidine 600 mg IV 30 min before the taxane.

Only immediate type hypersensitivity drug reactions involving IgE antibodies and/or a mast cell-mediated mechanism are considered eligible for desensitization. Administering increasing amounts of drug in an incremental and stepwise manner can provide a safe and effective means of continuing vital therapy while avoiding, or largely preventing, the adverse reactions. For carboplatin hypersensitivity, Lee et al. [250] administered a cumulative dose of 38.935 mg of drug in 11, 15-min steps followed by a final infusion over 184.4 min administering 461.065 mg (92 %). The total final cumulative dose of 500 mg was given over a 6–8 h period. In a rapid desensitization protocol for carboplatin, oxaliplatin, docetaxel and paclitaxel, Gastaminza et al. [240] employed five delivery stages in procedures lasting from 2 to 5 h with 89 % of the total dose delivered in 1 h in the last step. Once again, apart from data on the platins and taxanes, there is little pertinent information on desensitization for most other anti-cancer drugs. A successful desensitization protocol for the receptor tyrosine kinase inhibitor sunitinib involved ten escalating doses starting with 0.05 mg following premedication with prednisone and promethazine [251]. Examples of desensitization to imatinib have also been reported. Successes were achieved with gradual dose escalation over 10 weeks and, in another case, over 5 days starting with a dose of 0.005 mg and building up to a total of 400 mg [252]. In a 4-h outpatient oral procedure beginning with a 10-ng dose of imatinib, the drug was administered in increasing dosage every 15 min to ten patients, only two of whom ultimately proved unable to continue therapy [253].

9 The most insidious of all adverse consequences of targeted therapies: tumour adaptation, increased malignancy and the invasive metastatic switch

Over the last 15 years, molecularly targeted approaches involving innovative anti-angiogenic strategies, particularly drugs inhibiting the VEGF-A pathway, for example, the mAb bevacizumab and VEGFR tyrosine kinase inhibitors sorafenib and sunitinib [254], have brought about a revolution in the treatment of cancer. However, it has become increasingly clear that the complexities of tumour-induced angiogenesis and our understanding of its many interrelated processes and opposing effects on tumour resistance, growth and metastasis [255, 256] have produced outcomes that are often disappointing and more modest than originally expected [257]. From studies in tumour-bearing mice, the realization that anti-angiogenic therapy that initially exerts an antitumour
effect by inhibiting vascularization may also lead on to the opposing effects of enhanced tumour growth, invasiveness and distal metastasis has brought into focus the questions of how tumour resistance develops and how it can be overcome [257, 258]. A number of possible explanations have been advanced such as the involvement of pro-angiogenic cytokines and growth factors, e.g. granulocyte colony-stimulating factor (G-CSF), stromal cell-derived factor 1α (SDF-1α; CXCL12), osteopontin [259]; absence of VEGF or VEGFRs in metastatic tumours; tumours utilizing blood vessels in vascular-rich tissues; selection and growth of tumour cells resistant to hypoxia and a number of other suggestions [258, 260]. Tumour blood vessels are abnormal [261], and poor oxygenation is a well-known feature of solid tumours leading to tumour invasiveness that is due, in part at least, to increased expression of hypoxia-inducible factor-1α (HIF-1α) [262]. Several studies have highlighted the importance of improving oxygenation of cancer cells by demonstrating differentiation, vascular normalization and reduced metastasization following pO2 increases [263, 264] after promotion of the so-called invasive metastatic switch [265] in a hypoxic tumour environment [266, 267]. In findings that seem to offer one of the most promising early strategies to overcome the invasive and metastatic effects seen during resistance to anti-angiogenic drugs, the endogenous anti-angiogenic chemorepulsive protein semaphorin 3A normalized the vasculature and blocked tumour growth when re-expressed in mouse pancreatic neuroendocrine tumours and cervical carcinomas [268, 269]. HIF-1α which regulates cellular adaptation to oxygen deprivation is often associated with treatment failure [267], and its expression in human tumours has been shown to correlate with metastasis [270]. Treatment of tumour-bearing mice with sunitinib and semaphorin 3A demonstrated a marked reduction in HIF-1α levels that were previously elevated in animals treated with the anti-angiogenesis drug. Administration of semaphorin 3A together with sunitinib also reduced NF-κB and counteracted sunitinib-induced activation of c-Met (Met tyrosine kinase receptor; hepatocyte growth factor (HGF) receptor) and other signalling pathways active in hypoxia [269, 271]. NF-κB activates HIF-1α and cancer invasiveness and may have an important role in the development of resistance to anti-angiogenic resistance [272]. C-Met and its ligand, HGF (scatter factor) are upregulated in a number of different cancers where they contribute to angiogenesis and tumour growth and metastasis [273].

10 Combination therapies with targeted agents: adverse effects as a consequence of overcoming resistance to anti-angiogenic therapy

As summarized above, resistance eventually develops to VEGF inhibition therapy with the occurrence of residual primary tumour growth and/or metastasis [256]. It seems that restoration of the vasculature is the forerunner of oxygenation of tumour tissue and vascular normalization is critical not only for overcoming resistance to anti-angiogenic therapy but also to achieve success with such therapy together with other chemotherapeutic agents. These findings have ramifications for future targeted therapeutic approaches with anti-angiogenic agents and for consequent adverse events occurring in treated patients. HIF-1α is overexpressed in many cancers and its role in angiogenesis, tumour growth and metastasis is firmly established [266, 267] making the transcription factor a priority target for drug development [274, 275]. Topotecan, a topoisomerase I inhibitor that inhibits re-ligation of DNA strand breaks (Table 2), independently inhibits HIF-1α accumulation in human cancer cells [276] while the related drug irinotecan (Table 2) has already been employed in combination with bevacizumab to inhibit HIF-1α in recurrent glioblastoma [277]. Other non-targeted chemotherapeutic drugs that inhibit HIF-1α include the aryl hydrocarbon receptor aminoflavone, the peptide antibiotic with anticancer activity echinomycin and anthracyclines such as doxorubicin [278]. Targeted drugs that inhibit HIF-1α include HDAC and mTOR (everolimus and temsirolimus) inhibitors and the proteasome inhibitor bortezomib (Table 3) [279]. The mTOR inhibitor rapamycin together with irinotecan proved active in a mouse model of human metastatic colon cancer. The drug combination produced a reduction in tumour volume together with cell death under hypoxic but not normoxic conditions indicating a cytotoxic effect mediated via HIF-1α [280]. Inhibitors that target other pathways, particularly pathways to tumour invasiveness, are also being sought. One such pathway is the HGF/C-Met pathway where preclinical studies with inhibitors that block multiple receptor tyrosine kinases (including VEGF) and c-Met showed amplification of the effects of VEGFR blockade leading to reduced angiogenesis, increased apoptosis and reductions in tumour growth and metastasis [281].

For metastatic carcinoma, e.g. RCC, approved targeted therapies that have become available in recent years for first- and second-line use include sorafenib, sunitinib, pazopanib (Table 3), the VEGF-targeted mAb bevacizumab and mTOR inhibitors everolimus and temsirolimus (Table 3). The main adverse events to bevacizumab comprise GI perforation, haemorrhage, wound healing complications, thrombosis, infusion reactions, hypertension, necrotizing fasciitis, proteinuria and pulmonary events [282]; reactions to the other five drugs are summarized in Table 3. Axitinib (Inlyta®) which inhibits multiple receptor tyrosine kinases including VEGFRs, PDGFR and c-KIT was approved by the FDA in January 2012 for patients with advanced RCC after failure of one prior systemic therapy. Adverse reactions to the drug are quite extensive and include hypertension, thrombotic events, haemorrhage, GI perforation and fistula, hypothyroidism,
wound healing complications, proteinuria, reversible posterior leukoencephalopathy syndrome, hepatic impairment, hand-foot skin reaction and exfoliative dermatitis [283]. In addition to the well-known adverse reactions to the drugs used for mono-anti-angiogenic treatment, for example sunitinib or sorafenib, the likely increasing use of combination therapies to overcome the severe adverse consequences of the resistance-promoting targeted anti-angiogenic therapies will see the appearance of adverse responses to cytotoxic, non-targeted drugs like topotecan, irinotecan and anthracyclines with their well-known myelosuppression, cytopenias and occasional type I (anaphylactic) and types II and III hypersensitivity reactions (Table 2). As well as the employment of targeted mTOR, HDAC (e.g. vorinostat) or proteasome (e.g. bortezomib, carfilzomib) inhibitors will give rise to the known systemic and cutaneous reactions evoked by these drugs (Table 3). Drugs that inhibit hypoxia signalling are already being sought and examined [284], and it is certain that each new drug will provoke its own spectrum of adverse effects. Suggested new therapeutic approaches, for example employment of semaphorin A3 [269], to inhibit angiogenesis while assisting the development of the surviving vasculature and thus avoiding ongoing hypoxia are likely to be intensively examined to see whether the observed effects in mice also occur in man. Here it will be necessary to determine if the encouraging effects of inhibition of Met activation, increase in levels of the apparent tumour suppressor and adhesion molecule E-cadherin and reductions in tumour invasiveness and metastasis are not opposed by processes promoting tumour progression. The complexity of the tumour environment with its myriad effector pathways demands investigation of the latter possibility.

Other underlying mechanisms with the promise of possible new therapies currently being examined or under development include a soluble decoy receptor for VEGF [VEGF trap; aflibercept (Zaltrap®)]—for adverse reactions see Table 3 [285], inhibition of VEGF and angiopoietins [286], drugs that inhibit placental growth factor (PIGF) [287], targeting of angiopoietin-Tie2 signalling [288], the possible need to add anti-inflammatory therapy as a result of an inflammatory environment helping to sustain tumour progression [289], the search for inhibitors of matrix-degrading enzymes [e.g. matrix metalloproteinase (MMP)-9] for use as blockers of tumour invasion [290] and study of vascular integrin inhibitors for anti-angiogenic therapy. In relation to this last strategy, cilengitide, a cyclic RGD (Arg-Gly-Asp)-motif-containing pentapeptide (Fig. 6) that binds with high specificity to the αvβ3 and αvβ5 vascular integrin heterodimers, targets both tumour cells and the vasculature [291]. Integrins, a family of cell adhesion transmembrane receptors, may be highly expressed in some tumours; for example, αvβ3 and αvβ5 are significantly upregulated in human gliomas. Cilengitide, the first integrin inhibitor to reach phase III clinical development, shows promise for glioblastoma with patients demonstrating good tolerance to what are claimed to be the drug’s relatively few side effects. The most common adverse events reported to the FDA during the period 1997–2012 were effects on the white blood cells, viz. neutropenia and thrombocytopenia as well as pulmonary embolism, urinary tract infections, anorexia, joint and bone pain and electrolyte imbalance. A recent announcement from the phase III CENTRIC trial which investigated the efficacy and tolerability of cilengitide in newly diagnosed glioblastoma patients with methylated O(6)-methylguanine DNA methyltransferase gene promoter status revealed that the drug did not meet its primary endpoint of significantly increasing overall survival.

It is clear that many currently favoured strategies to elucidate mechanisms and find new and effective anti-cancer agents are leading to targeted ‘multi-modal’ therapies combining different anti-angiogenic, or anti-angiogenic together with cytotoxic, approaches [292]. It should be anticipated that as the number and variety of therapeutic agents given to a patient increases (and these may ultimately include some unusual choices, e.g. anti-inflammatory drugs), the problem of sorting out and quickly identifying the drug(s) responsible for an adverse reaction inevitably becomes more difficult.

11 Concluding remarks: do some hypersensitivity responses go unrecognised? Towards multi-modal treatment strategies

A consideration of the possibility of hypersensitivity reactions is a necessary part of clinical drug trials, and such reactions are commonly referred to in information released by drug regulatory agencies, drug package inserts, the oncology literature, some pharmacological investigations and toxicology studies. The term hypersensitivity is widely used across a number of disciplines, but it is clear that those using the term do not always have the same definition in mind. Even within its ‘home’ disciplines of immunology and allergy, there is inconsistency and confusion over both the definition of hypersensitivity and the assignment of the term, and this situation has not been helped by the WAO’s so-called ‘stringent definition’ based, as claimed, on ‘the mechanisms that initiate and mediate allergic reactions’ without including any reference to allergic and/or immune involvement [7]. With such a definition, it is difficult to see how types I to IV hypersensitivity reactions, distinguished on the basis of humoral and cellular immune mediation, can be separated from a variety of other reactions showing ‘objectively reproducible symptoms or signs’ initiated by ‘a defined stimulus at a dose tolerated by normal persons’. It is not surprising therefore to find that a drug reaction classified as a hypersensitivity response in the absence of any test investigations might be anything from an
unsurprising collection of respiratory or gastrointestinal symptoms to a full-blown bullous skin eruption. Such inconsistent use of the term hypersensitivity is particularly apparent in the assessment and recording of adverse reactions to chemotherapeutic drugs. Patients sometimes show different and unexpected reactions to the same drug. The question of the possible participation of humoral or cellular immune mechanisms in anti-neoplastic drug-induced thrombocytopenia, neutropenia, anaemia, vascular disorders, liver injury and lung disease as well as in the many dermatological manifestations is obviously fundamental in any attempted assessment of the incidence of true hypersensitivity responses to the increasing battery of drugs used in large numbers of patients for therapy of many different cancers. Even a quick perusal of Tables 2 and 3 reveals that adverse haematological, liver, lung and skin reactions occur across the multitude of chemically diverse drugs, many with vastly different modes of action, but it is frequently not at all clear whether a reaction is a true hypersensitivity response and if any immune process, direct or indirect, is involved. In a recent review of immune haemolytic anaemia associated with drug therapy, Garraty [293] summarized the current situation in words that apply as much to drug-induced reactions to platelets, neutrophils, the vasculature, liver, lung and cutaneous cells as to red cells: ‘With increased knowledge of the immune system… it is a pity that more modern approaches have not been applied to solving the mechanism(s) involved in drug-induced RBC auto antibody formation’.

If widespread acceptance of what constitutes a hypersensitivity reaction is to be achieved, agreement on the definition of such reactions must occur and that will require the acceptance of uniform criteria for some common terminology. In the standard classification and severity grading scale in cancer therapy, clinical trials and other oncology settings published as Common Terminology Criteria for Adverse Events by the US National Institutes of Health and NCI under the US Cancer Therapy Evaluation Program, ‘hypersensitivity’ is grouped with ‘allergic reaction’ [234], in itself indicating the acceptance of an immune basis for hypersensitivity responses. In recent years, progress has been made in elucidating mechanisms underlying a range of drug-induced cell-mediated type IV cutaneous eruptions (e.g. allergic contact dermatitis, psoriasis, erythema multiforme, AGEP, SJS, TEN) and systemic/cutaneous reactions (e.g. DRESS) [210], and together with our understanding of the mechanisms of types I, II and III hypersensitivities, it should be possible to present clear
guidelines on the criteria necessary to identify a reaction as a true hypersensitivity response. In everyday practice, it will sometimes be difficult to clearly establish an immune basis for a reaction since, as outlined above, suitable tests necessary to define the mechanism of the reaction may not be available or easily accessible. Obtaining the necessary evidence to be sure of a hypersensitivity classification for a particular reaction may be especially difficult in some cases of suspected drug-induced immune lung disease, liver injury and vascular disorders where work needs to be done to develop and apply suitable tests. Even for drug-induced thrombocytopenia and neutropenia, proving immune mediation of reactions will be difficult for some investigators, clinicians and facilities. However, despite the admittedly inadequate testing procedures available, it is likely that systematic application of the existing skin test approaches and in vitro assays could significantly improve the discrimination of many haematological, vascular, lung, liver and skin reactions and identify some true types I, II, III and IV drug hypersensitivities that may currently go unidentified because of the present widespread absence of any such testing. Existing assays and procedures for detecting drug-specific/reactive IgE and cytotoxic antibodies, drug-antibody immune complexes and drug-reactive T cells need to be more frequently employed, but further research is needed to improve the range, specificity, ease of use and ready access of procedures for the identification of the four types of hypersensitivities to the many different anti-cancer drugs now so widely used. This area of oncoimmunology research has been neglected, not least by immunologists and allergists, to the detriment of patients, our recognition of the true nature and spectrum of side effects of many anti-neoplastic drugs and our understanding of some important and still poorly researched vascular, lung, liver and cutaneous drug-induced reactions.

Clearly, the terminology used to describe hypersensitivity reactions needs to be standardised, but while the unsatisfactory lack of a common definition of what constitutes a hypersensitivity reaction and hence the resultant misuse of the term are particularly apparent in cancer chemotherapy, the same deficiencies, perhaps to a somewhat lesser extent, also exist in the allergy field. Greater involvement of both clinical and experimental allergists in the assessment of adverse drug responses to neoplastic agents in oncology would be of benefit to both disciplines and, importantly, lead to a more accurate interpretation of adverse events during clinical trials of cancer drugs. Adverse reactions that are serious enough to threaten further administration of drugs vital for therapy of particular cancers occur commonly and fatal outcomes may result in cases of drug withdrawal when this might have otherwise been prevented by skilled desensitization procedures. Given this, the potential value of skin testing in some cases and the need for premedication treatments and desensitizations, it is both surprising and unfortunate that allergists and clinical immunologists often seem to be so little involved in the management of adverse/hypersensitivity reactions in cancer chemotherapy.

Finally, in relation to the immediate future of cancer chemotherapy, understanding the basis of the resistance that develops to VEGF inhibition therapy by, for example receptor tyrosine kinase inhibitors such as sorafenib and sunitinib, has brought with it the realization of the need to restore normal vasculature and oxygenation of tumour tissue. Arising from this is the desire to identify some fresh therapeutic targets such as the transcription factor HIF-1α, the expression of which correlates with metastasis. Multi-modal treatment strategies combining anti-angiogenesis and cytotoxic therapies will see the appearance of already known side effects of the ‘older’ cytotoxic drugs topotecan, irinotecan and some anthracyclines, some existing targeted drugs including mTOR, HDAC and proteasome inhibitors but, as well, the yet to be determined reactions to the new therapeutic strategies under investigation/development such as those targeting angiopoietin-Tie2 signalling, semaphorins for angiogenesis inhibition and inhibitors of HIF-1α, c-Met, E-cadherin, VEGF-angiopoietins, PIGF, MMPs and vascular integrins.

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