Metformin and cancer: Quo vadis et cui bono?

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ABSTRACT

How many lives have already been saved by the anti-cancer drug metformin? Inadvertently perhaps, among the millions of type 2 diabetics with occult or known cancers and who have been prescribed metformin since the 1950s, thousands may have benefitted from the anticancer properties of this first-line pharmacotherapy. Quo vadis? Now, researchers aim to move metformin from a non-targeted stage of cancer therapy that has been mostly developed retrospectively and empirically into a targeted therapy by following a biological rationale and a predefined mechanism of action. But, who might benefit from metformin? Cui bono? Because metformin is on the leading edge of a new generation of cancer metabolism-targeted therapies, perhaps it is the right time to provide solutions to the challenges that metformin and other onco-biguanides will face in the coming years before becoming incorporated into the therapeutic armamentarium against cancer.

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In a highly prevalent disease such as breast cancer (BC), a moderate improvement in survival rates would be enough to save thousands of lives worldwide if a given therapeutic approach is widely available and is accompanied by a benefit to a significant proportion of women developing the disease. When considering insulin as one of the most plausible biological mechanisms underlying the link between obesity and diabetes and a significantly increased risk of BC mortality, it is remarkable that a lowering of insulin by 25% is associated with a 5% absolute improvement in BC mortality [1]. Given that metformin can reduce fasting insulin levels 10-30% in women with early BC [2], one could speculate that metformin may have saved thousands of lives among the millions of diabetic women receiving metformin in the last 60 years. To put these putative numbers into a relevant scenario, we should acknowledge that a shift in the treatment paradigm of adjuvant tamoxifen treatment from the more traditional 5 years to 10 years after diagnosis of BC was praised as “life saving” by the mainstream media when the landmark ATLAS study [3] demonstrated the cumulative risk of dying of BC between 5 and 14 years after first diagnosis was 12.2% in the extended treatment group vs 15% in the standard treatment group, a reduction in absolute risk of 2.8% (or a reduction in BC mortality of 28 per 1,000 women). If risk of breast (and other) cancer is significantly increased in the protracted period when hyperinsulinemia is present before diabetes diagnosis, as well as the strong association between hyperinsulinemia and the metabolic syndrome, countless lives might be saved because more than 25% of the world’s adult population would benefit from metformin’s ability to reduce the insulin-related risk of cancer and cancer-related mortality. However, while cancer researchers and patients are eagerly awaiting the results of the NCIC Clinical Trials Group MA.32 study—an ongoing adjuvant trial of 3,640 women with early stage BC examining the effect of metformin vs placebo on invasive cancer-free survival and...
other outcomes with completed surgery and (neo)adjuvant chemotherapy [4]-there is growing uncertainty regarding the best models to predict the practical metformin posology that might be required for clinical activity in nondiabetic subjects.

On the one hand, numerous translational concerns have been expressed over the growing list of in vitro studies that are oftently claimed clinically irrelevant because millimolar concentrations of metformin are typically required to induce anticancer effects [5-8], whereas the doses used in the clinical management of type 2 diabetes are normally in the range of 8-30 µM. However, the so-called metformin dosing paradox might not be translationally significant when considering that a dynamic relationship exists between metformin efficacy and chronic energy excess [9-12]. Because glucose availability promotes cancer aggressiveness while reducing metformin efficacy, maintenance of glucose homeostasis through energy restriction, exercise, and other pharmacological modalities is expected to enhance metformin responsiveness and clinical outcomes in cancer patients [11]. Alternatively, synthetic lethality could be exploited to overcome glucose-driven resistance to metformin by combining metformin with anti-angiogenesis agents that promote severe glucose deprivation in certain areas of the tumor [12].

Table 1: Summary of metformin posology in murine cancer models

| Study/Year          | Animal strain | Route of administration | Plasma concentration (mean, [range]) |
|---------------------|---------------|-------------------------|-------------------------------------|
| Memmott et al. (2010) | A/J           | Oral (Dose/time)        | i.p. (Dose/time)                    |
|                     |               | 1000 mg/kg/13 weeks    | 250 mg/kg/13 weeks                  | 13 µM                          |
|                     |               |                        |                                    | 31 µM                          |
| Menendez et al. (2014) | BALB/c nu/nu | 250 mg/kg/8 weeks      | 200 mg/kg/8 weeks                  | 5 µM [3-6]                      |
|                     |               |                        |                                    | 674 µM [387-1161]              |
| Chandel et al. (2016) | C57BL/6J     | 250 mg/kg/2 weeks      | -                                   | 5 µM                           |
|                     |               |                        |                                    | -                              |
| Chandel et al. (2016) | J:Nu          | 250 mg/kg/2 weeks      | -                                   | [3.2-12.4 µM]                  |
|                     |               |                        |                                    | -                              |
| Chandel et al. (2016) | NMRI nu/nu   | -                      | 350 mg/kg/2 weeks                  | -                              |
|                     |               |                        |                                    | 7.5 µM                         |
| Dowling et al. (2016) | NOD/SCID     | -                      | 125 mg/kg/0.5 hr                   | -                              |
|                     |               |                        |                                    | 184 µM [61-288]                |
| Dowling et al. (2016) | NOD/SCID     | 1000 mg/kg/16 days     | 125 mg/kg/15 days                  | 34 µM [2-126]                  |
|                     |               |                        |                                    | 145 µM [66-215]                |

and its implications for human therapy [13, 14].

The studies led by Dowling and Chandel recapitulate earlier findings from our laboratory and others showing that mice i.p. injected with metformin have considerably higher plasma levels of metformin than those receiving the drug dissolved in the drinking water [15, 16] (Table 1). To evaluate the missing PK properties of metformin in the context of cancer therapy while concurrently excluding a role for insulin in its anti-tumor action in murine cancer models, we used breast xenotumors formed by PIK3CA H1047R-mutated MCF10DCIS.com cells (Cufí et al., 2013), which proliferate in vitro irrespective of the presence or absence of insulin, and form tumors that are refractory to dietary restriction (DR) in vivo [17, 18]. In a first cohort, cell line-derived xenograft (CDX) mice had ad libitum access to drinking water with 1.25 mg/mL metformin, which provided a daily oral dose of ~250 mg/kg. In a second cohort, metformin was administered to CDX mice by i.p. injection (200 mg/kg, once daily). The average plasma levels of metformin were evaluated using HPLC coupled to electrospray ionization quadrupole time-of-flight mass spectrometry at the end of an 8-week treatment when animals were sacrificed to evaluate tumor volume, mitotic activity, and anatomopathological features. BALB/c nude mice receiving metformin in the drinking water yielded plasma concentrations in the 5 µM range, an identical range to that reported by Chandel et al. [14] in C57NL/6J mice and in J:Nu mice bearing the metformin-responsive HCT116 p53- colon cancer tumor (Table 1). When metformin was measured 2 h after the last i.p. dose, metformin peaked in the plasma ~100 fold higher than that found with the oral dosing schedule. Dowling et al. [13] now similarly report that the average plasma level of metformin in NOD/SCID mice xenografted with metformin-unresponsive HCT116 colorectal cancer cells was drastically higher in the i.p.-treated group (Table 1). Because the cation metformin is expected to accumulate up to 500-fold in mitochondria by virtue of the membrane potential, the low micromolar concentrations of plasma
metformin in murine cancer models treated with the standard oral dosing of diabetic patients might translate into an anti-tumor activity exclusively attributable to metformin’s ability to inhibit oxidative phosphorylation [13]. In our hands, such diabeto-biguanide activity modestly affected the growth of insulin-independent breast xenotumors, reaching a maximum of 43% at 4 weeks after inoculation of tumor cells and decreasing to 30-35% reduction toward the end of the treatment period [16, 18]. The apparent correlation between longer exposure to i.p.-delivered metformin, higher levels of plasma metformin, and higher efficacy in murine cancer models might suggest a re-evaluation of metformin administration regimens to optimize drug plasma levels and delivery to the tumor, but with the potential of involving mechanisms of action not strictly related with blockade of oxidative phosphorylation [14]. In our hands, such onco-biguanide activity increased in a temporal manner, reaching a highly significant >80% inhibition of tumor growth [16, 18].

We need to anticipate the challenges that developing metformin and other onco-biguanides as bona fide anti-cancer metabolic therapeutics will face in the coming years (Table 2):

- First, the recent discovery that in vivo environment strictly dictates the metabolic phenotype of tumors in patients and mouse models [19] together with the contradictory findings obtained in patient-derived xenograft (PDX) models treated with metformin [20, 21], strongly indicate that clinically-translatable models remain to be developed to accurately assess the true potential of metformin-based therapeutic modalities. The use of immunocompromised mice or the fact that the subcutaneous compartment where tumors are placed in CDX and PDX is not analogous to the microenvironment within organs might significantly impact the response to metformin. Patient-derived orthotopic xenografts (PDOX) and 3D cancer tissue bioreactors for long-term organotypic culture and drug delivery might better approximate a metformin-targetable in vivo tumor microenvironment ex vivo.

- Second, effective design, analysis, and interpretation of PK and pharmacodynamic (PD) studies would enable a better understanding of the mechanism of metformin action and identify PK properties for further improvement. A valuable route to implement the successful development of metformin as an anti-cancer drug might involve PK/PD modeling of metformin in type 2 diabetics enrolled in early research phases of other oncology discovery projects involving metformin-related molecular targets (e.g., inhibitors of mTOR, PI3K) [22-24]. Such an approach could help elucidate the relationship between PK and PD, help to understand the mechanism of metformin action, and identify PK properties for further development and optimal usage of metformin in a cancer setting.

- Third, a major challenge in metformin posology and/or administration to optimize drug plasma levels is how to ensure that high metformin concentrations will cause tumor regression with minimal normal tissue toxicity. Although it is commonly acknowledged that such approaches will inevitably suffer from narrow therapeutic windows that might cause side effects such as lactic acidosis, we must consider the evidence that a Cochrane review and an analysis of the UK-based General Practice Research Database revealed no cases of fatal lactic acidosis and crude incidence rates of only 3.3 cases per 100,000 person-years among metformin users [25, 26]. Although it could be argued that additional side effects might be seen at supra-diabetic metformin concentrations, there have been no attempts to evaluate metformin-based synthetic lethal interactions sparing normal cells while selectively killing cancer cells. Hypothesis-driven and screening-based (e.g., chemical, siRNA/shRNA, CRISPR libraries) synthetic lethality approaches [27] would identify second-site molecular targets that could synergize with metformin-related metabolic targets and improve metformin efficacy, and might even provide new clinically relevant biomarkers for patient selection. A hypothesis-driven synthetic lethality approach recently conducted in our laboratory revealed the extreme vulnerability of cells deficient in the BC early onset gene BRCA1 to NAD+ depletion after concurrent metformin [28] and nicotinamide phosphoribosyltransferase (NAMPT) inhibition, suggesting that the evaluation of metformin/

| CHALLENGE | PROPOSAL |
|-----------|----------|
| (1) Clinically translatable models | CDX*, PDX® → PDOX®, 3-D tissue bioreactors |
| (2) Implementation of PK/PD strategies | Metformin PK/PD in type 2 diabetics being enrolled in early research phases of oncology drugs targeting metformin-related molecular targets |
| (3) Therapeutic window and patient selection | Identification of synthetically lethal interactions to improve metformin-induced selective killing of cancer cells while sparing normal cells |
| (4) Companion diagnostics | Identification of metabolomic/fluxomic signatures that can be used both to predict efficacy and safety (outcome) and to monitor the response to metformin |

*aCDX: Cell line-derived xenografts; bPDX: Patient-derived xenografts; bPDOX: Patient-derived orthotopic xenografts
NAMPT small molecule inhibitor combinations (e.g., FK866) to treat BRCA1-related BC may be warranted (manuscript in preparation).

- Fourth, identification of predictive biomarkers of response to metformin and precise patient selection is essential to ensure the measurement of true response rates without bias through inclusion of patients who do not respond to metformin simply because they are not suitable candidates to benefit from it. In the vast majority of ongoing metformin-based clinical trials, patient selection based on a particular biomarker has not been carried out, which may have improved the commonly observed poor response rates. In addition, the dysregulation and integration of multiple metabolic pathways in most tumors might trigger redundant/compensatory networks that could impact the effectiveness of metformin. Metformin-diagnostic co-development programs should be implemented to identify “decisive” stratification factors that have the potential to be an important tool for clinicians in relation to: 1) the identification of patients who are most likely to benefit from metformin; 2) the identification of patients likely to be at increased risk of serious adverse reactions as a result of metformin treatment; and 3) monitoring response to metformin to achieve improved safety or effectiveness. Such companion diagnostic assays, which should have a high degree of analytical validity before they can be released for routine clinical usage, might incorporate multi-metabolite panels based on the identification of metformin-driven metabolomic/fluxomic “fingerprints” of specific cellular events related with efficacy and safety. The exploration of the exo-metabolome to monitor, in real-time, biomarker/surrogate endpoints of metformin efficacy in liquid biopsies including urine [29] might optimize and accelerate the design of metformin-based personalized cancer therapies.

Inadvertently perhaps, among the millions of type 2 diabetics with occult or known cancers and who have been prescribed metformin since the 1950s, thousands may have benefited from the anticancer properties of this first-line pharmacotherapy. Quo vadis? Now, researchers aim to move metformin from a “non-targeted” stage of cancer therapy that has been mostly developed retrospectively and empirically into a “targeted” therapy by following a biological rationale and a predefined mechanism of action. But, who might benefit from metformin? Cui bono? If metformin is on the leading edge of a new generation of metabolic targeted therapies, perhaps it is the right time to consider that the challenges facing the incorporation of metformin into the therapeutic armamentarium against cancer might be as complicated as targeting genetic aberrations, if not more so.

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CONFLICTS OF INTEREST

The authors have no financial conflicts of interest.

Author contributions

J.A.M, B. M-C., and J.J. developed the study concept. J.A.M, B. M-C., and J.J. interpreted data. B. M-C., is the principal investigator of the neoadjuvant metformin trial METTEN-01 (https://www.clinicaltrialsregister.eu/ctr-search/search?query=2011-000490-30). J.A.M wrote the paper. All authors approved the manuscript for publication.

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